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EP 0 936 507 A2

(12)

EUROPEAN PATENT APPLICATION

(43) Date of publication: 18.08.1999 Bulletin 1999/33

(51) Int Cl.⁶: **G03G 9/083**, G03G 9/09, C01G 49/00

(21) Application number: 99301117.0

(22) Date of filing: 16.02.1999

(84) Designated Contracting States:
AT BE CH CY DE DK ES FI FR GB GR IE IT LI LU
MC NL PT SE
Designated Extension States:
AL LT LV MK RO SI

(30) Priority: 17.02.1998 JP 5282698

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(54) Black magnetic composite particles and black magnetic toner using the same

(57) Black magnetic composite particles for a black magnetic toner comprising:

(a) magnetic iron oxide particles having an average particle diameter of from 0.055 to 0.95 μ m; (b) a coating on the surface of said magnetic iron oxide particles, comprising at least one organosilicon compound selected from:

(1) organosilane compounds obtainable by dry-

ing or heat-treating alkoxysilane compounds, (2) polysiloxanes or modified polysiloxanes, and

(3) fluoroalkyl organosilane compounds obtainable by drying or heat-treating fluoroalkylsilane compounds; and

(c) carbon black fine particles adhered on at least a part of said coating, which have a particle size of from 0.002 to 0.05 μ m and which are present in an amount of from 1 to 25 parts by weight per 100 parts by weight of said magnetic iron oxide particles.

Descripti n

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[0001] Th present invention relates to black magnetic composit particles and a black magnetic toner, and more particularly, to black magnetic composite particles for black magnetic toner which not only show an exc llent disp rsibility in a binder resin due to less amount of carbon black fine particles fallen-off from the surface of each particle, but also have an excellent fluidity and blackness, and a black magnetic toner using such black magnetic composite particles

[0002] As one of conventional electrostatic latent image-developing methods, there has been widely known and generally adopted a so-called one component system development method of using as a developer, a magnetic toner comprising composite particles prepared by mixing and dispersing magnetic particles such as magnetite particles in a resin, without using a carrier.

[0003] The conventional development methods of using one-component magnetic toner have been classified into CPC development methods of using a low-resistance magnetic toner, and PPC development methods of using a high-resistance magnetic toner.

[0004] In the CPC methods, the low-resistance magnetic toner used therefor has an electric conductivity, and is charged by the electrostatic induction due to electric charge of the latent images. However, since the charge induced on the magnetic toner is lost while the magnetic toner is transported from a developing zone to a transfer zone, the low-resistance magnetic toner is unsuitable for the PPC development method of using an electrostatic transfer method. In order to solve this problem, there have been developed the insulated or high resistance magnetic toners having a volume resistivity as high as not less than 10¹⁴ Ω -cm.

[0005] As to the insulated or high-resistance magnetic toner, it is known that the developing characteristics thereof are affected by magnetic particles exposed to the surface of the magnetic toner, or the like.

[0006] Recently, with the high image quality such as high image density or high tone gradation, or with the high copying speed of duplicating machines, it has been strongly demanded to further enhance characteristics of the insulted or high-resistance magnetic toners as a developer, especially a fluidity thereof.

[0007] With respect to such demands, in Japanese Patent Application Laid-Open (KOKAI) No. 53-94932(1978), there has been described "these high-resistance magnetic toners are deteriorated in fluidity due to the high electric resistance, so that there arises such a problem that non-uniformity of developed images tend to be caused. Namely, although the high-resistance magnetic toners for PPC development method can maintain necessary charges required for image transfer, the magnetic toners are frictionally charged even when they are present in other steps than the transfer step, where the magnetic toners are not required to be charged, e.g., in a toner bottle or on the surface of a magnetic roll, or also slightly charged by mechano-electrets during the production process of these magnetic toners. Therefore, the magnetic toners tend to be electrostatically agglomerated, resulting in deterioration of fluidity thereof", and "It is an another object of the present invention to provide a high-resistance magnetic toner for PPC development method which is improved in fluidity, can be prevented from causing non-uniformity of developed images, and has an excellent image definition and tone gradation, thereby obtaining high-quality copies by indirect copying methods".

[0008] In recent years, with the reduction in particle size of the insulated or high-resistance magnetic toners, it has been increasingly required to enhance the fluidity thereof.

[0009] With respect to such a fact, in "Comprehensive Data Collection for Development and Utilization of Toner Materials" published by Japan Scientific Information Co., Ltd., page 121, there has been described "With extensive development of printers such as ICP, a high image quality has been required. In particular, it has been demanded to develop high-precision or high-definition printers. In Table 1, there is shown a relationship between definitions obtained by using the respective toners. As is apparent from Table 1, the smaller the particle size of wet toners, the higher the image definition is obtained. Therefore, when a dry toner is used, in order to enhance the image definition, it is also required to reduce the particle size of the toner ···· As reports of using toners having a small particle size, it has been proposed that by using toners having a particle size of 8.5 to 11 μm, fogs on a background can be improved and toner consumption can be reduced, and further by using polyester-based toners having a particle size of 6 to 10 μm, an image quality, a charging stability and lifetime of the developer can be improved. However, when such toners having a small particle size are used, it has been required to solve many problems. There are problems such as improvement in productivity, sharpness of particle size distribution, improvement in fluidity, etc.".

[0010] Further, black magnetic toners widely used at the present time, have been required to show a high degree of blackness and a high image density for line images and solid area images on copies.

[0011] With respect to this fact, on page 272 of the above-mentioned "Compreh nsiv Data Collection for Development and Utilization of Toner Materials", there has been described "Powder development is characterized by a high image density. How v r, th high image density as w II as th fog d nsity as described her inafter, greatly influences image characteristics obtained".

[0012] Ther is a close r lationship betw in propinties of the magnetic ton r and thos of the magnetic particles mixed and dispersed in thi magnetic toner.

[0013] That is, the fluidity of the magnetic ton r is largely varied depending upon surface condition of the magnetic particles exposed to the surface of the magnetic toner. Therefore, the magnetic particles sthems the largely required to show an excellent fluidity.

[0014] The degree of blackness and density of the magnetic toner are also largely varied depending upon the degree of blackness and density of the magnetic particles as a black pigment contained in the magnetic toner.

[0015] As the black pigment, magnetite particles have been widely used from the standpoints of magnetic properties such as saturation magnetization or coercive force, low price, color tone or the like. In addition to the magnetite particles, carbon black fine particles may be added. However, in the case where the carbon black fine particles are used in a large amount, the electric resistance is lowered, so that it is not possible to obtain an insulated or high-resistance magnetic toner.

[0016] Hitherto, in order to enhance the fluidity of the black magnetic toner, there have been many attempts of improving the fluidity of the magnetite particles mixed and dispersed in the magnetic toner. For example, there have been proposed (1) a method of forming spherical-shaped magnetite particles (Japanese Patent Application Laid-Open (KOKAI) No. 59-64852(1984)), (2) a method of exposing a silicon compound to the surface of magnetite particles (Japanese Patent Publication (KOKOKU) No. 8-25747(1996)), or the like.

[0017] Black magnetic particles for black magnetic toner, which have not only an excellent fluidity and blackness, but also an excellent dispersibility in a binder resin, are presently strongly demanded. However, black magnetic particles capable of satisfying all of these requirements have not been obtained yet.

[0018] Namely, the above-mentioned spherical magnetite particles show a higher fluidity than those of cubic magnetite particles, octahedral magnetite particles or the like. However, the fluidity of the spherical magnetite particles is still insufficient, and further the blackness is disadvantageously low.

[0019] As to the above-mentioned magnetite particles to the surface of which the silicon compound is exposed, the fluidity thereof is also still insufficient, and the blackness thereof is also disadvantageously low.

[0020] As a result of the present inventor's earnest studies for solving the above problems, it has been found that by coating magnetic iron oxide particles having an average particle size of 0.055 to 0.95 μm with an organosilicon compound, then adhering carbon black fine particles having an average particle size of 0.002 to 0.05 μm on a coating of the organosilicon compound to obtain black magnetic composite particles, a black magnetic toner produced by using the obtained black magnetic composite particles, can have not only an excellent fluidity and an excellent blackness, but also can show a high-resistance or an insulating property without lowering in the electric resistance. The present invention has been attained on the basis of the finding.

[0021] It is an object of the present invention to provide black magnetic particles for black magnetic toner, which are not only excellent in fluidity and blackness, but also can show an excellent dispersibility in a binder resin.

[0022] It is another object of the present invention to provide a black magnetic toner, which is not only excellent in fluidity and blackness, but also small in reduction of electric resistance and, therefore, can realize a high image quality and a high copying speed.

[0023] To accomplish the aims, in a first aspect of the present invention, there is provided black magnetic composite particles for a black magnetic toner, comprising:

magnetic iron oxide particles having an average particle diameter of 0.055 to 0.95 μ m; a coating formed on surface of said magnetic iron oxide particles, comprising at least one organosilicon compound selected from the group consisting of:

- (1) organosilane compounds obtained by drying or heat-treating an alkoxysilane compounds,
- (2) polysiloxanes or modified polysiloxanes, and

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(3) fluoroalkyl organosilane compounds obtained by drying or heat-treating a fluoroalkylsilane compounds; and

carbon black fine particles adhered on at least a part of said coating, having a particle size of 0.002 to $0.05\,\mu m$, the amount of said carbon black fine particles adhered being 1 to 25 parts by weight based on 100 parts by weight of said magnetic iron oxide particles.

[0024] In a second aspect of the present invention, there is provided black magnetic composite particles for a black magnetic toner, comprising:

magnetic iron oxide particles having an average particle diameter of 0.055 to $0.95 \,\mu m$, and having a coat which is formed on at least a part of the surfac of said magnitic iron oxide particles and which comprises at least one compound sell cted from the group consisting of hydroxides of aluminum, oxides of aluminum, hydroxid is of silicon and oxides of silicon in an amount of 0.01 to $50 \,\%$ by we light, calculated as Al or SiO_2 , based on the total weight of the magnetic iron oxide particles;

a coating formed on surface of said magnetic iron oxide particles, comprising at I ast one organosilicon compound selected from the group consisting of:

- (1) organosilane compounds obtained by drying or heat-treating an alkoxysilane compounds,
- (2) polysiloxanes or modified polysiloxanes, and
- (3) fluoroalkyl organosilane compounds obtained by drying or heat-treating a fluoroalkylsilane compounds; and

carbon black fine particles adhered on at least a part of said coating, having a particle size of 0.002 to 0.05 μm, the amount of said carbon black fine particles adhered being 1 to 25 parts by weight based on 100 parts by weight of said magnetic iron oxide particles.

[0025] In a third aspect of the present invention, there is provided black magnetic toner comprising:

said black magnetic composite particles comprising: magnetic iron oxide particles having an average particle diameter of 0.055 to 0.95 µm; a coating formed on surface of said magnetic iron oxide particles, comprising at least one organosilicon compound selected from the group consisting of:

- (1) organosilane compounds obtained by drying or heat-treating an alkoxysilane compounds,
- (2) polysiloxanes or modified polysiloxanes, and
- (3) fluoroalkyl organosilane compounds obtained by drying or heat-treating a fluoroalkylsilane compounds; and

carbon black fine particles adhered on at least a part of said coating, having a particle size of 0.002 to 0.05 μ m, the amount of said carbon black fine particles adhered being 1 to 25 parts by weight based on 100 parts by weight of said magnetic iron oxide particles, and a binder resin.

[0026] In a fourth aspect of the present invention, there is provided black magnetic toner comprising:

30 said black magnetic composite particles comprising:

magnetic iron oxide particles having an average particle diameter of 0.055 to 0.95 μ m, and having a coat which is formed on at least a part of the surface of said magnetic iron oxide particles and which comprises at least one compound selected from the group consisting of hydroxides of aluminum, oxides of aluminum, hydroxides of silicon and oxides of silicon in an amount of 0.01 to 50 % by weight, calculated as Al or SiO_2 , based on the total weight of the magnetic iron oxide particles;

a coating formed on surface of said magnetic iron oxide particles, comprising at least one organosilicon compound selected from the group consisting of:

- (1) organosilane compounds obtained by drying or heat-treating an alkoxysilane compounds,
- (2) polysiloxanes or modified polysiloxanes, and
- (3) fluoroalkyl organosilane compounds obtained by drying or heat-treating a fluoroalkylsilane compounds; and

carbon black fine particles adhered on at least a part of said coating, having a particle size of 0.002 to 0.05 μ m, the amount of said carbon black fine particles adhered being 1 to 25 parts by weight based on 100 parts by weight of said magnetic iron oxide particles, and a binder resin.

[0027] In a fifth aspect of the present invention, there is provided a method of using black magnetic composite particles for production of a black magnetic toner, which black magnetic composite particles comprise comprising:

magnetic iron oxide particles having an average particle diameter of 0.055 to $0.95 \,\mu m$; a coating formed on surface of said magnetic iron oxide particles, comprising at least one organosilicon compound selected from the group consisting of:

- (1) organosilane compounds obtain d by drying or heat-tr ating an alkoxysilan compounds,
- (2) polysiloxan s or modified polysiloxan s, and
- (3) fluoroalkyl organosilane compounds obtain d by drying or heat-treating a fluoroalkylsilane compounds; and

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carbon black fine particles adhered on at least a part of said coating, having a particle size of 0.002 to $0.05 \, \mu m$, the amount of said carbon black fin particles adhered being 1 to 25 parts by weight based on 100 parts by weight of said magnetic iron oxide particles.

5 In the accompanying drawings

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[0028] Fig. 1 is an electron micrograph (× 20,000) showing a particle structure of spherical magnetite particles used in Example 1.

[0029] Fig. 2 is an electron micrograph (× 20,000) showing a particle structure of carbon black particles used in Example 1.

[0030] Fig. 3 is an electron micrograph (× 20,000) showing a particle structure of black magnetic composite particles obtained in Example 1.

[0031] Fig. 4 is an electron micrograph (× 20,000) showing a particle structure of mixed particles composed of the spherical magnetite particles and the carbon black fine particles, for comparative purpose.

[0032] The present invention is now described in detail below.

[0033] First, the black magnetic composite particles according to the present invention are described.

[0034] The black magnetic composite particles according to the present invention, comprise magnetic iron oxide particles as core particles having an average particle diameter of 0.055 to $095\,\mu\text{m}$, a coating comprising an organosilicon compound which is formed on the surface of each magnetic iron oxide particle, and carbon black fine particles having an average particle size of 0.002 to $0.05\,\mu\text{m}$, adhered on at least a part of the coating comprising the organosilicon compound.

[0035] As the magnetic iron oxide particles used as core particles in the present invention, there may be exemplified magnetite particles ($\text{FeO}_x \text{-Fe}_2\text{O}_3$; $0 < \text{X} \le 1$), maghemite particles ($\gamma \text{-Fe}_2\text{O}_3$) or a mixture of these particles. In the consideration of blackness of the obtained black magnetic composite particles, magnetite particles are preferred.

[0036] As the magnetic iron oxide particles as core particles, from the viewpoint of a particle shape thereof, there may be exemplified isotropic particles having a ratio of an average particle length (average major diameter) to an average particle breadth (average minor diameter) of usually not less than 1.0 and less than 2.0, preferably 1.0 to 1.8, more preferably 1.0 to 1.5, such as spherical particles, granular particles or polyhedral particles, e.g., hexahedral particles or octahedral particles, or anisotropic particles having an aspect ratio (average major axial diameter/average minor axial diameter; hereinafter referred to merely as "aspect ratio") of not less than 2:1, such as acicular particles, spindle-shaped particles or rice ball-shaped particles. In the consideration of the fluidity of the obtained black magnetic composite particles, the magnetic iron oxide particles having an isotropic shape are preferred. Among them, the spherical particles are more preferred.

[0037] In the case of the isotropic magnetic iron oxide particles, the average particle size (diameter) thereof is 0.055 to 0.95 μ m, preferably 0.065 to 0.75 μ m, more preferably 0.065 to 0.45 μ m. In the case of the anisotropic magnetic iron oxide particles, the average major axial diameter thereof is 0.055 to 0.95 μ m, preferably 0.065 to 0.75 μ m, more preferably 0.065 to 0.45 μ m, and the aspect ratio (average major axial diameter/average minor axial diameter) thereof is 2:1 to 20:1, preferably 2:1 to 15:1, more preferably 2:1 to 10:1.

[0038] When the average particle size of the magnetic iron oxide particles is more than $0.95\,\mu m$, the obtained black magnetic composite particles are coarse particles and are deteriorated in tinting strength. On the other hand, when the average particle size is less than $0.055\,\mu m$, the intermolecular force between the particles is increased due to the reduction in particle size (fine particle), so that agglomeration of the particles tends to be caused. As a result, it becomes difficult to uniformly coat the surfaces of the magnetic iron oxide particles with the organosilicon compounds, and uniformly adhere the carbon black fine particles on the surface of the coating comprising the organosilicon compounds.

[0039] Further, in the case where the upper limit of the aspect ratio of the anisotropic magnetic iron oxide particles exceeds 20:1, the particles tend to be entangled with each other, and it also becomes difficult to uniformly coat the surfaces of the magnetic iron oxide particles with the organosilicon compounds, and uniformly adhere the carbon black fine particles on the surface of the coating composed of the organosilicon compounds.

[0040] As to the particle size distribution of the magnetic iron oxide particles, the geometrical standard deviation value thereof is preferably not more than 2.0, more preferably not more than 1.8, still more preferably not more than 1.6. When the geometrical standard deviation value thereof is more than 2.0, coarse particles are contained therein, so that the particles are inhibited from being uniformly dispersed. As a result, it also becomes difficult to uniformly coat the surfaces of the magnetic iron oxide particles with the organisilicon compounds, and uniformly adhere the carbon black fine particles on the surface of the coating composed of the organosilicon compounds. The lower limit of the geometrical standard deviation value is 1.01. It is industrially difficult to obtain particles having a geometrical standard deviation value of less than 1.01.

[0041] The BET specific surfac area of the magn tic iron oxide particles there f is not less than 0.5 m²/g. Wh n the BET specific surfac area is I ss than 0.5 m²/g, the magn tic iron oxide particles may become coars particles,

or the sintering between the particles may be caused, so that the obtained black magnetic composit particles also may become coarse particles and tind to be deteriorated in tinting strength. In the consideration of the tinting strength of the obtain displayed black magnetic composite particles, the BET specific surface area of the magnetic iron oxide particles is preferably not less than 1.0 m²/g, more preferably 3.0 m²/g. Further, in the consideration of uniformly coating the surfaces of the magnetic iron oxide particles with the organosilicon compounds, and uniformly adhering the carbon black fine particles on a coating composed of the organosilicon compounds, the upper limit of the BET specific surface area of the magnetic iron oxide particles, is usually 70 m²/g, preferably 50 m²/g, more preferably 20 m²/g.

[0042] As to the fluidity of the magnetic iron oxide particles, the fluidity index thereof is about 25 to about 44. Among the magnetic iron oxide particles having various shapes, the spherical particles are excellent in fluidity, for example, the fluidity index thereof is about 30 to about 44.

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[0043] As to the blackness of the magnetic iron oxide particles, in the case of the magnetite particles, the lower limit thereof is usually 18.0 when represented by L* value, and the upper limit thereof is usually 25.0, preferably 24.0 when represented by L* value. In the case of maghemite particles, the lower limit thereof is usually more than 18.0 when represented by L* value, and the upper limit thereof is usually 32, preferably 30 when represented by L* value. When the L* value exceeds the above-mentioned upper limit, the lightness of the particles is increased, so that it is difficult to obtain black magnetic composite particles having a sufficient blackness.

[0044] As to the magnetic properties of the magnetic iron oxide particles, the coercive force value thereof is usually about 10 to about 350 Oe, preferably 20 to about 330 Oe; the saturation magnetization value in a magnetic field of 10 kOe is usually about 50 to about 91 emu/g, preferably about 60 to about 90 emu/g; and the residual magnetization value in a magnetic field of 10 kOe is usually about 1 to about 35 emu/g, preferably about 3 to about 30 emu/g.

[0045] The particle shape and particle size of the black magnetic composite particles according to the present invention are considerably varied depending upon those of the magnetic iron oxide particles as core particles. The black magnetic composite particles have a similar particle shape to that of the magnetic iron oxide particle as core particle, and a slightly larger particle size than that of the magnetic iron oxide particles as core particles.

[0046] More specifically, when the isotropic magnetic iron oxide particles are used as core particles, the obtained black magnetic composite particles according to the present invention, have an average particle size of usually 0.06 to 1.0 μ m, preferably 0.07 to 0.8 μ m, more preferably 0.07 to 0.5 μ m and a ratio of an average particle length to an average particle breadth of usually not less than 1.0 and less than 2.0, preferably 1.0 to 1.8, more preferably 1.0 to 1.5,. When the anisotropic magnetic iron oxide particles are used as core particles, the obtained black magnetic composite particles according to the present invention, have an average particle size of usually 0.06 to 1.0 μ m, preferably 0.07 to 0.8 μ m, more preferably 0.07 to 0.5 μ m.

[0047] When the average particle size of the black magnetic composite particles is more than 1.0 µm, the obtained black magnetic composite particles may be coarse particles, and deteriorated in tinting strength. On the other hand, when the average particle size thereof is less than 0.06 µm, the black magnetic composite particles tends to be agglomerated by the increase of intermolecular force due to the reduction in particle size, thereby deteriorating the dispersibility in a binder resin upon production of the magnetic toner.

[0048] When the anisotropic magnetic iron oxide particles are used as core particles, the upper limit of the aspect ratio of the obtained black magnetic composite particles according to the present invention, is usually 20:1, preferably 18:1, more preferably 15:1. When the aspect ratio is more than 20:1, the black magnetic composite particles may be entangled with each other in the binder resin, so that the dispersibility in binder resin tends to be deteriorated.

[0049] The geometrical standard deviation value of the black magnetic composite particles according to the present invention is preferably not more than 2.0, more preferably 1.01 to 1.8, still more preferably 1.01 to 1.6. The lower limit of the geometrical standard deviation value thereof is preferably 1.01. When the geometrical standard deviation value thereof is more than 2.0, the tinting strength of the black magnetic composite particles is likely to be deteriorated due to the existence of coarse particles therein. It is industrially difficult to obtain such particles having a geometrical standard deviation of less than 1.01.

[0050] The BET specific surface area of the black magnetic composite particles according to the present invention, is usually 1 to 200 m²/g, preferably 2 to 150 m²/g, more preferably 2.5 to 100 m²/g. When the BET specific surface area thereof is less than 1 m²/g, the obtained black magnetic composite particles may be coarse, and the sintering between the black magnetic composite particles is caused, thereby deteriorating the tinting strength. On the other hand, when the BET specific surface area is more than 200 m²/g, the black magnetic composite particles tend to be agglomerated together by the increase in intermolecular force due to the reduction in particle size, thereby deteriorating the dispersibility in a bindires in upon production of the magnetic toner.

[0051] As to the fluidity of the black magnetic composite particles according to the present invention, the fluidity index thereof is preferably 45 to 80, more preferably 46 to 80, still more preferably 47 to 80. When the fluidity index thereof is less than 45, the fluidity of the black magnetic composite particles becomes insufficient, the reby failing to improve the fluidity of the finally obtained magnetic toner. Further, in the production process of the magnetic toner, the retend to be caused defects such as clogging of hopper, etc., the reby distribution of the production process of the magnetic toner.

[0052] As to the blackness of the black magnetic composite particles according to the present invintion, in the case magnetite particles are used as core particles, the upper limit of thi blackniss of the black magnetic composite particles is usually 20.0, preferably 19.0, more preferably 18.0 when right presented by L* value. In this case maghemite particles are used as core particles, the upper limit of the blackness of the black magnetic composite particles is usually 20.0, preferably 19.5, more preferably 19.0 when represented by L* value. When the L* value thereof is more than 20.0, the lightness of the obtained black magnetic composite particles becomes high, so that the black magnetic composite particles having a sufficient blackness cannot be obtained. The lower limit of the blackness thereof is 15 when represented by L* value.

[0053] The dispersibility in binder resin of the black magnetic composite particles according to the present invention, is preferably 4th or 5th rank, more preferably 5th rank when evaluated by the method described hereinafter.

[0054] The percentage of desorption of carbon black fine particles from the black magnetic composite particles according to the present invention, is preferably not more than 20 %, more preferably not more than 10 %. When the desorption percentage of the carbon black fine particles is more than 20 %, the desorbed carbon black fine particles tend to inhibit the black magnetic composite particles from being uniformly dispersed in the binder resin upon production of the magnetic toner.

[0055] The magnetic properties of the black magnetic composite particles according to the present invention, can be controlled by appropriately selecting kind and particle shape of the magnetic iron oxide particles as core particles. Similarly to magnetic properties of magnetic particles ordinarily used for the production of magnetic toner, the coercive force of the black magnetic composite particles according to the present invention, is usually about 10 to about 350 Oe, preferably about 20 to about 330 Oe; the saturation magnetization in a magnetic field of 10 kOe is usually about 50 to about 91 emu/g, preferably about 60 to about 90 emu/g; and the residual magnetization in a magnetic field of 10 kOe is usually about 1 to about 35 emu/g, preferably about 3 to about 30 emu/g.

[0056] The coating formed on the surfaces of the core particles comprises at least one organosilicon compound selected from the group consisting of (1) organosilane compounds obtained by drying or heat-treating alkoxysilane compounds; (2) polysiloxanes, or modified polysiloxanes selected from the group consisting of (A) polysiloxanes modified with at least one compound selected from the group consisting of polyethers, polyesters and epoxy compounds (hereinafter referred to merely as "modified polysiloxanes"), and (B) polysiloxanes whose molecular terminal is modified with at least one group selected from the group consisting of carboxylic acid groups, alcohol groups and a hydroxyl group; and (3) fluoroalkyl organosilane compounds obtained by drying or heat-treating fluoroalkylsilane compounds.

[0057] The organosilane compounds (1) can be produced by drying or heat-treating alkoxysilane compounds represented by the formula (I):

$$H_a^1 SiX_{4-a}$$
 (I)

wherein R¹ is C_6H_6 -, $(CH_3)_2CHCH_2$ - or $n-C_bH_{2b+1}$ - (wherein b is an integer of 1 to 18); X is CH_3O - or C_2H_6O -; and a is an integer of 0 to 3.

[0058] The drying or heat-treatment of the alkoxysilane compounds is conducted, for example, at a temperature of usually 40 to 200°C, preferably 60 to 150°C for usually 10 minutes to 12 hours, preferably 30 minutes to 3 hours.

[0059] Specific examples of the alkoxysilane compounds may include methyl triethoxysilane, dimethyl diethoxysilane, tetraethoxysilane, phenyl triethyoxysilane, diphenyl diethoxysilane, methyl trimethoxysilane, dimethyl dimethoxysilane, tetramethoxysilane, phenyl trimethoxysilane, diphenyl dimethoxysilane, isobutyl trimethoxysilane, decyl trimethoxysilane or the like. Among these alkoxysilane compounds, in view of the desorption percentage and the adhering effect of carbon black fine particles, methyl triethoxysilane, phenyl triethyoxysilane, methyl trimethoxysilane, dimethyl dimethoxysilane and isobutyl trimethoxysilane are preferred, and methyl triethoxysilane and methyl trimethoxysilane are more preferred.

[0060] As the polysiloxanes (2), there may be used those compounds represented by the formula (II):

$$\begin{array}{cccc}
CH_3 & R^2 & CH_3 \\
CH_3 - Si - O & - Si - O \\
CH_3 & CH_3 & CH_3
\end{array}$$
(II)

wherein R^2 is H- or CH_3 -, and d is an int g r of 15 to 450.

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[0061] Among thes polysiloxanes, in view of the desorption percentage and the adhering effect of carbon black fine particles, polysiloxanes having methyl hydrogen siloxane units ar pr ferr d.

[0062] As th modified polysiloxanes (A), there may be used:

(a) polysiloxanes modified with polyethers represented by the formula (III):

wherein \mathbb{R}^3 is -(- $\mathbb{C}H_2$ -)_n-; \mathbb{R}^4 is -(- $\mathbb{C}H_2$ -)_r- $\mathbb{C}H_3$; \mathbb{R}^5 is -OH, -COOH, - $\mathbb{C}H_2$ -CH₂, - $\mathbb{C}(\mathbb{C}H_3$)= $\mathbb{C}H_2$ or-(- $\mathbb{C}H_2$ -)_r- $\mathbb{C}H_3$; \mathbb{R}^5 is -(- $\mathbb{C}H_2$ -)_k- $\mathbb{C}H_3$; $\mathbb{C}H_3$; $\mathbb{C}H_3$; $\mathbb{C}H_3$ and $\mathbb{C}H_3$ is an integer of 1 to 15; $\mathbb{C}H_3$; $\mathbb{C}H_3$; $\mathbb{C}H_3$ is an integer of 1 to 300;

(b) polysiloxanes modified with polyesters represented by the formula (IV):

wherein R^7 , R^8 and R^9 are -(- CH_2 -)_q- and may be the same or different; R^{10} is -OH, -COOH, - CH_2 -CH₂, - $C(CH_3)$ = CH_2 or -(- CH_2 -)_r- CH_3 ; R^{11} is-(- CH_2 -)_e- CH_3 ; n and q are an integer of 1 to 15; r and s are an integer of 0 to 15; e' is an integer of 1 to 50; and f' is an integer of 1 to 300;

(c) polysiloxanes modified with epoxy compounds represented by the formula (V):

wherein H^{12} is -(-CH₂-)_v-; v is an integer of 1 to 15; t is an integer of 1 to 50; and u is an integer of 1 to 300; or a mixture thereof.

[0063] Among these modified polysiloxanes (A), in view of the desorption percentage and the adhering effect of carbon black fin particles, the polysiloxanes modified with the polyether represented by the formula (III), are preferred.

[0064] As the terminal-modified polysiloxanes (B), there may be used those represented by the formula (VI):

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wherein R¹³ and R¹⁴ are -OH, R¹⁶OH or R¹⁷COOH and may be the same or different; R¹⁵ is -CH₃ or -C₆H₅; R¹⁶ and R¹⁷ are -(-CH₂-)_v-; y is an integer of 1 to 15; w is an integer of 1 to 200; and x is an integer of 0 to 100.

[0065] Among these terminal-modified polysiloxanes, in view of the desorption percentage and the adhering effect of carbon black fine particles, the polysiloxanes whose terminals are modified with carboxylic acid groups are preferred.

[0066] The fluoroalkyl organosilane compounds (3) may be produced by drying or heat-treating fluoroalkylsilane compounds represented by the formula (VII):

$$CF_3(CF_2)_zCH_2CH_2(H^{18})_a,SiX_{4-a'}$$
 (VII)

wherein H_3^{18} is CH_3 -, C_2H_5 -, CH_3O - or C_2H_5O -; X is CH_3O - or C_2H_5O -, and z is an integer of 0 to 15; and a' is an integer of 0 to 3.

[0067] The drying or the heat-treatment of the fluoroalkylsilane compounds may be conducted, for example, at a temperature of usually 40 to 200°C, preferably 60 to 150°C for usually 10 minutes to 12 hours, preferably 30 minutes to 3 hours.

[0068] Specific examples of the fluoroalkylsilane compounds may include trifluoropropyl trimethoxysilane, tridecaftuoroctyl trimethoxysilane, heptadecafluorodecyl trimethoxysilane, heptadecafluorodecyl triethoxysilane, trifluoropropyl triethoxysilane, tridecafluoroctyl triethoxysilane, heptadecafluorodecyl triethoxysilane, heptadecafluorodecyl triethoxysilane, heptadecafluorodecyl triethoxysilane, heptadecafluorodecyl triethoxysilane or the like. Among these fluoroalkylsilane compounds, in view of the desorption percentage and the adhering effect of carbon black fine particles, trifluoropropyl trimethoxysilane, tridecafluoroctyl trimethoxysilane and heptadecafluorodecyl trimethoxysilane are preferred, and trifluoropropyl trimethoxysilane and tridecafluoroctyl trimethoxysilane are more preferred.

[0069] The coating amount of the organosilicon compounds is usually 0.02 to 5.0 % by weight, preferably 0.03 to 2.0 % by weight, more preferably 0.05 to 1.5 % by weight (calculated as Si) based on the weight of the acicular hematite particles or the acicular iron oxide hydroxide particles coated with the organosilicon compounds.

[0070] When the coating amount of the organosilicon compounds is less than 0.02 % by weight, it becomes difficult to adhere the carbon black fine particles on the surfaces of the magnetic iron oxide particles in such an amount enough to improve the fluidity and blackness of the obtained black magnetic composite particles.

[0071] On the other hand, when the coating amount of the organosilicon compounds is more than 5.0 % by weight, a sufficient amount of the carbon black fine particles can be adhered on the surfaces of the magnetic iron oxide particles. However, the use of such unnecessarily large amount of the organosilicon compounds is meaningless because the effect of enhancing the fluidity or blackness of the obtained black magnetic composite particles is already saturated.

[0072] As the carbon black fine particles used in the black magnetic composite particles according to the present invention, there may be exemplified commercially available carbon black particles such as furnace black, channel black or the like. Specific examples of the commercially available carbon black particles usable in the present invention, may include MA100, MA7, #1000, #2400B, #30, MA8, MA11, #50, #52, #45, #2200B, MA600, etc. (tradenames; produced by Mitsubishi Chemical Corp.), Seast9H, Seast7H, Seast6, Seast3H, Seast300, SeastFM, etc. (tradenames; produced by Tokai Carbon Co., Ltd.) or the like. In the consideration of compatibility with the organosilicon compounds, MA100, MA7, #1000, #2400B and #30 are preferred.

[0073] The average particle size of the carbon black fine particles is usually about 0.002 to about 0.035 μ m, preferably about 0.002 to about 0.035 μ m.

50 [0074] When the average particle size of the carbon black fine particles is less than 0.002 μm, the carbon black fine particles are too fine to be well handled.

[0075] On the other hand, when the average particle size of the carbon black fine particles is more than 0.05 µm, the particle size of the carbon black fine particle second seco

[0076] The ratio of the average particle size of the magnetic iron oxide particles to that of the carbon black fine particles is preferably not less than 2:1. When the ratio is less than 2:1, the particle size of the carbon black fine particles becomes considerably larger as compared to that of the magnetic iron oxide particles as core particles, thereby causing insufficient adhesion of the carbon black fine particles onto the coating composed of the organosilicon compounds, and increasing the desorption percentage of the carbon black fine particles. As a result, the obtained black magnetic composite particles tend to be deteriorated in dispersibility in a binder resin upon the production of magnetic toner.

[0077] The amount of the carbon black fine particles adhered is 1 to 25 parts by weight based on 100 parts by weight of the magnetic iron oxide particles as core particles.

[0078] When the amount of the carbon black fine particles adhered is less than 1 part by weight, the amount of the carbon black fine particles adhered is insufficient, so that it becomes difficult to obtain black magnetic composite particles having a sufficient fluidity and blackness.

[0079] On the other hand, when the amount of the carbon black fine particles adhered is more than 25 parts by weight, the obtained black magnetic composite particles can show a sufficient fluidity and blackness. However, since the amount of the carbon black fine particles adhered is considerably large, the carbon black fine particles tend to be desorbed from the coating composed of the organosilicon compound. As a result, the obtained black magnetic composite particles tend to be deteriorated in dispersibility in a binder resin upon the production of magnetic toner.

[0080] In the black magnetic composite particles according to the present invention, the surfaces of the magnetic iron oxide particles as core particles may be preliminarily coated with at least one compound selected from the group consisting of hydroxides of aluminum, oxides of aluminum, hydroxides of silicon and oxides of silicon (hereinafter referred to as "coating composed of hydroxides and/or oxides of aluminum and/or silicon"), if necessary. In this case, the obtained black magnetic composite particles can show a higher dispersibility in a binder resin as compared to in the case where the magnetic iron oxide particles are uncoated with hydroxides and/or oxides of aluminum and/or silicon.

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[0081] The coating amount of the hydroxides and/or oxides of aluminum and/or silicon is preferably 0.01 to 50 % by weight (calculated as Al, SiO₂ or a sum of Al and SiO₂) based on the weight of the magnetic iron oxide particles as core particles.

[0082] When the coating amount of the hydroxides and/or oxides of aluminum and/or silicon is less than 0.01 % by weight, the effect of enhancing the dispersibility of the obtained black magnetic composite particles in a binder resin upon the production of magnetic toner cannot be obtained.

[0083] On the other hand, when the coating amount of the hydroxides and/or oxides of aluminum and/or silicon is more than 50 % by weight, the obtained black magnetic composite particles can exhibit a good dispersibility in a binder resin upon the production of magnetic toner. However, the use of such unnecessarily large coating amount of the hydroxides and/or oxides of aluminum and/or silicon is meaningless.

[0084] The particle size, geometrical standard deviation, BET specific surface area, fluidity, blackness L* value and desorption percentage of carbon black fine particles of the black magnetic composite particles wherein the surface of the core particle is coated with the hydroxides and/or oxides of aluminum and/or silicon according to the present invention, are substantially the same as those of the black magnetic composite particles wherein the core particle is uncoated with the hydroxides and/or oxides of aluminum and/or silicon according to the present invention.

[0085] The black magnetic composite particles according to the present invention can be produced by the following method.

[0086] Among the isotropic magnetite particles which are magnetic iron oxide particles, (1) octahedral magnetite particles can be produced by passing an oxygen-containing gas through a suspension containing ferrous hydroxide colloid having a pH value of not less than 10, which is obtained by reacting an aqueous ferrous salt solution with an aqueous alkali solution having a concentration of not less than one equivalent based on Fe2+ in the aqueous ferrous salt solution, thereby precipitating magnetite particles, and then subjecting the obtained magnetite particles to filtering, washing with water and drying (Japanese Patent Publication (KOKOKU) No. 44-668(1969); (2) hexahedral magnetite particles can be produced by passing an oxygen-containing gas through a suspension containing ferrous hydroxide colloid having a pH value of 6.0 to 7.5, which is obtained by reacting an aqueous ferrous salt solution with an aqueous alkali solution having a concentration of not more than one equivalent based on Fe^{2+} in the aqueous ferrous salt solution to produce magnetite core particles, further passing an oxygen-containing gas through the obtained aqueous ferrous salt reaction solution containing the magnetite core particles and the ferrous hydroxide colloid, at a pH value of 8.0 to 9.5, to precipitate magnetite particles, and then subjecting the precipitated magnetite particles to filtering, washing with water and drying (Japanese Patent Application Laid-Open (KOKAI) No. 3-201509(1991); (3) spherical magnetite particles can be produced by passing an oxygen-containing gas through a suspension containing ferrous hydroxide colloid having a pH value of 6.0 to 7.5, which is obtained by reacting an aqui ous ferrous salt solution with an aqueous alkali solution having a conc ntration of not more than on equival nt bas d on Fe2+ in the aqueous ferrous salt solution to produce magn title core particles, adding alkali hydroxide in an amount of not less than equivalent based on the remaining Fe²⁺ to adjust the pH value of the suspension to not less than 10, heat-oxidizing the resultant suspension to precipitate magnetite particles, and then subjecting the precipitated magnetit particles to filtering, washing with water

and drying (Japanese Patent Publication (KOKOKU) No. 62-51208(1987).

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[0087] The isotropic maghemite particles can be obtained by heating the above-mention of isotropic magnetic titles in air at 300 to 600°C.

[0088] The anisotropic magnetite particles can be produced by passing an oxygen-containing gas through a suspension containing either ferrous hydroxide colloid, iron carbonate, or an iron-containing precipitate obtained by reacting an aqueous ferrous salt solution with alkali hydroxide and/or alkali carbonate, while appropriately controlling the pH value and temperature of the suspension, to produce acicular, spindle-shaped or rice ball-shaped goethite particles, subjecting the obtained goethite particles to filtering, washing with water and drying, and then reducing the goethite particles in a heat-reducing gas at 300 to 800°C.

[0089] The anisotropic maghemite particles can be produced by heat-oxidizing the above-mentioned anisotropic magnetite particles in an oxygen-containing gas at 300 to 600°C.

[0090] The coating of the magnetic iron oxide particles with the alkoxysilane compounds, the polysiloxanes, the modified polysiloxanes or the fluoroalkylsilane compounds, may be conducted by mechanically mixing and stirring the magnetic iron oxide particles together with the alkoxysilane compounds, the polysiloxanes, the modified polysiloxanes, the terminal-modified polysiloxanes or the fluoroalkylsilane compounds; or by mechanically mixing and stirring both the components together while spraying the alkoxysilane compounds, the polysiloxanes, the modified polysiloxanes, the terminal-modified polysiloxanes or the fluoroalkylsilane compounds onto the magnetic iron oxide particles. In these cases, substantially whole amount of the alkoxysilane compounds, the polysiloxanes, the modified polysiloxanes, the terminal-modified polysiloxanes or the fluoroalkylsilane compounds added can be applied onto the surfaces of the magnetic iron oxide particles.

[0091] In order to uniformly coat the surfaces of the magnetic iron oxide particles with the alkoxysilane compounds, the polysiloxanes, the modified polysiloxanes, the terminal-modified polysiloxanes or the fluoroalkylsilane compounds, it is preferred that the magnetic iron oxide particles are preliminarily diaggregated by using a pulverizer. As the apparatuses for the mixing and stirring, there may be used an edge runner, a Henschel mixer or the like.

[0092] The conditions for the mixing and stirring such as mixing ratio, linear load, stirring speed or mixing and stirring time, may be appropriately adjusted so as to coat the surfaces of the magnetic iron oxide particles with the alkoxysilane compounds, the polysiloxanes, the modified polysiloxanes, the terminal-modified polysiloxanes or the fluoroalkylsilane compounds as uniformly as possible. The mixing and stirring time for the coating treatment is, for example, preferably not less than 20 minutes.

[0093] The amount of the alkoxysilane compounds, the polysiloxanes, the modified polysiloxanes, the terminal-modified polysiloxanes or the fluoroalkylsilane compounds added, is preferably 0.15 to 45 parts by weight based on 100 parts by weight of the magnetic iron oxide particles. When the amount of the organosilicon compounds added is less than 0.15 part by weight, it may become difficult to adhere the carbon black fine particles in such an amount enough to improve the blackness and flowability of the obtained black magnetic composite particles. On the other hand, when the amount of the organosilicon compounds added is more than 45 parts by weight, a sufficient amount of the carbon black fine particles can be adhered on the surface of the coating, but it is meaningless because the blackness and volume resistivity of the composite particles cannot be further improved by using such an excess amount of the organosilicon compounds.

[0094] Next, the carbon black fine particles are added to the magnetic iron oxide particles coated with the organosilicon compounds, and the resultant mixture is mixed and stirred to adhere the carbon black fine particles on the surfaces of the coating composed of the organosilicon compounds, and then dried.

[0095] In the case where the alkoxysilane compounds (1) and the fluoroalkylsilane compounds (3) are used as the coating compound, after the carbon black fine particles are adhered on the surface of the coating, the resultant composite particles are dried or heat-treated, for example, at a temperature of usually 40 to 200°C, preferably 60 to 150°C for usually 10 minutes to 12 hours, preferably 30 minutes to 3 hours, thereby forming a coating composed of the organosilane compounds (1) and the fluoroalkyl organosilane compounds (3), respectively.

[0096] It is preferred that the carbon black fine particles are added little by little and slowly, especially about 5 to 60 minutes.

[0097] The conditions for mixing and stirring the magnetic iron oxide particles and the carbon black fine particles coated with the organosilicon compounds, such as mixing ratio, linear load stirring speed or mixing and stirring time, may be appropriately adjusted so as to uniformly adhere the carbon black fine particles on the surface of the coating. The mixing and stirring time for the adhesion treatment is, for example, preferably not less than 20 minutes.

[0098] The amount of the carbon black fin particl s added, is preferably 1 to 30 parts by weight bas d on 100 parts by weight of the magnetic iron oxide particles. When the amount of the carbon black fine particles added is less than 1 part by weight, it may become difficult to adhere the carbon black fine particles in such an amount enough to improve the blackness and flowability of the obtained composite particles. On the other hand, when the amount of the carbon black fine particles added is more than 30 parts by weight, a sufficient blackness and flowability of the resultant composite particles can be obtained, but the carbon black fine particles tend to be desorbed from the surface of the coating

becaus of too large amount of the carbon black fine particles adhered, resulting in deteriorat d dispersibility in the binder resin upon the production of the magnetic toner.

[0099] The magnetic iron oxide particles may be coated with at least on compound selected from the group consisting of hydroxides of aluminum, oxides of aluminum, hydroxides of silicon and oxides of silicon, if required, prior to mixing and stirring with the alkoxysilane compounds, the polysiloxanes, the modified polysiloxanes, the terminal-modified polysiloxanes or the fluoroalkylsilane compounds.

[0100] The coating of the hydroxides and/or oxides of aluminum and/or silicon may be conducted by adding an aluminum compound, a silicon compound or both the compounds to a water suspension in which the magnetic iron oxide particles are dispersed, followed by mixing and stirring, and further adjusting the pH value of the suspension, if required, thereby coating the surfaces of the magnetic iron oxide particles with at least one compound selected from the group consisting of hydroxides of aluminum, oxides of aluminum, hydroxides of silicon and oxides of silicon. The thus obtained particles coated with the hydroxides and/or oxides of aluminum and/or silicon are then filtered out, washed with water, dried and pulverized. Further, the particles coated with the hydroxides and/or oxides of aluminum and/or silicon may be subjected to posttreatments such as deaeration treatment and compaction treatment, if required.

[0101] As the aluminum compounds, there may be exemplified aluminum salts such as aluminum acetate, aluminum sulfate, aluminum chloride or aluminum nitrate, alkali aluminates such as sodium aluminate, alumina sols or the like.

[0102] The amount of the aluminum compound added is 0.01 to 50 % by weight (calculated as AI) based on the weight of the magnetic iron oxide particles. When the amount of the aluminum compound added is less than 0.01 % by weight, it may be difficult to sufficiently coat the surfaces of the magnetic iron oxide particles with hydroxides and/ or oxides of aluminum, thereby failing to achieve the improvement of the dispersibility in the binder resin upon the production of the magnetic toner. On the other hand, when the amount of the aluminum compound added is more than 50 % by weight, the coating effect is saturated and, therefore, it is meaningless to add such an excess amount of the aluminum compound.

[0103] As the silicon compounds, there may be exemplified #3 water glass, sodium orthosilicate, sodium metasilicate, colloidal silica or the like.

[0104] The amount of the silicon compound added is 0.01 to 50 % by weight (calculated as SiO₂) based on the weight of the magnetic iron oxide particles. When the amount of the silicon compound added is less than 0.01 % by weight, it may be difficult to sufficiently coat the surfaces of the magnetic iron oxide particles with hydroxides and/or oxides of silicon, thereby failing to achieve the improvement of the dispersibility in the binder resin upon the production of the magnetic toner. On the other hand, when the amount of the silicon compound added is more than 50 % by weight, the coating effect is saturated and, therefore, it is meaningless to add such an excess amount of the silicon compound.

[0105] In the case where both the aluminum and silicon compounds are used in combination for the coating, the total amount of the aluminum and silicon compounds added is preferably 0.01 to 50 % by weight (calculated as a sum of Al and SiO₂) based on the weight of the magnetic iron oxide particles.

[0106] Next, the black magnetic toner according to the present invention is described.

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[0107] The black magnetic toner according to the present invention comprises the black magnetic composite particles, and a binder resin. The black magnetic toner may further contain a mold release agent, a colorant, a charge-controlling agent and other additives, if necessary.

[0108] The black magnetic toner according to the present invention has an average particle size of usually 3 to 15 μ m, preferably 5 to 12 μ m.

[0109] The amount of the binder resin used in the black magnetic toner is usually 50 to 900 parts by weight, preferably 50 to 400 parts by weight based on 100 parts by weight of the black magnetic composite particles.

[0110] As the binder resins, there may be used vinyl-based polymers, i.e., homopolymers or copolymers of vinyl-based monomers such as styrene, alkyl acrylates and alkyl methacrylates. As the styrene monomers, there may be exemplified styrene and substituted styrenes. As the alkyl acrylate monomers, there may be exemplified acrylic acid, methyl acrylate, butyl acrylate or the like.

[0111] It is preferred that the above copolymers contain styrene-based components in an amount of usually 50 to 95 % by weight.

[0112] In the binder resin used in the present invention, the above-mentioned vinyl-based polymers may be used in combination with polyester-based resins, epoxy-based resins, polyurethane-based resins or the like, if necessary.

[0113] As to the fluidity of the black magnetic toner according to the present invention, the fluidity index is usually 70 to 100, preferably 71 to 100, more preferably 72 to 100. When the fluidity index is less than 70, the black magnetic toner may not show a sufficient fluidity.

[0114] The blackness of the black magnetic toner according to the present invention is usually not more than 20, preferably not more than 19.5 when r presented by L* valu. When the blackness thereof is more than 20, the lightness of the black magnetic tener may be increased, resulting in insufficient blackness. The lower limit of the blackness of the black magnetic toner is usually about 15 when represented by L* valu.

[0115] The volume r sistivity of the black magn tic ton r according to the present invention, is usually not less than

 $1.0 \times 10^{13} \Omega$ -cm, pref rably not less than $3.0 \times 10^{13} \Omega$ -cm, more preferably not less than $5.0 \times 10^{13} \Omega$ -cm. When the volume resistivity is less than $1.0 \times 10^{13} \Omega$ -cm, the charge amount of the black magnetic toner tends to vary depending upon environmental conditions in which the toner is used, resulting in unstable properties of the black magnetic toner. The upper limit of the volume resistivity is $1.0 \times 10^{17} \Omega$ -cm.

[0116] As to the magnetic properties of the black magnetic toner according to the present invention, the coercive force thereof is usually 10 to 350 Oe, preferably 20 to 330 Oe; the saturation magnetization value in a magnetic field of 10 kOe is usually 10 to 85 emu/g, preferably 20 to 80 emu/g; the residual magnetization in a magnetic field of 10 kOe is usually 1 to 20 emu/g, preferably 2 to 15 emu/g; the saturation magnetization in a magnetic field of 1 kOe is usually 7.5 to 65 emu/g, preferably 10 to 60 emu/g; and the residual magnetization in a magnetic field of 1 kOe is usually 0.5 to 15 emu/g, preferably 1.0 to 13 emu/g.

[0117] The black magnetic toner according to the present invention may be produced by a known method of mixing and kneading a predetermined amount of a binder resin and a predetermined amount of the black magnetic composite particles together, and then pulverizing the mixed and kneaded material into particles. More specifically, the black magnetic composite particles and the binder resin are intimately mixed together with, if necessary, a mold release agent, a colorant, a charge-controlling agent or other additives by using a mixer. The obtained mixture is then melted and kneaded by a heating kneader so as to render the respective components compatible with each other, thereby dispersing the black magnetic composite particles therein. Successively, the molten mixture is cooled and solidified to obtain a resin mixture. The obtained resin mixture is then pulverized and classified, thereby producing a magnetic toner having an aimed particle size.

[0118] As the mixers, there may be used a Henschel mixer, a ball mill or the like. As the heating kneaders, there may be used a roll mill, a kneader, a twin-screw extruder or the like. The pulverization of the resin mixture may be conducted by using pulverizers such as a cutter mill, a jet mill or the like. The classification of the pulverized particles may be conducted by known methods such as air classification, etc., as described in Japanese Patent No. 2683142 or the like.

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[0119] As the other method of producing the black magnetic toner, there may be exemplified a suspension polymerization method or an emulsion polymerization method. In the suspension polymerization method, polymerizable monomers and the black magnetic composite particles are intimately mixed together with, if necessary, a colorant, a polymerization initiator, a cross-linking agent, a charge-controlling agent or the other additives and then the obtained mixture is dissolved and dispersed together so as to obtain a monomer composition. The obtained monomer composition is added to a water phase containing a suspension stabilizer while stirring, thereby granulating and polymerizing the composition to form magnetic toner particles having an aimed particle size.

[0120] In the emulsion polymerization method, the monomers and the black magnetic composite particles are dispersed in water together with, if necessary, a colorant, a polymerization initiator or the like and then the obtained dispersion is polymerized while adding an emulsifier thereto, thereby producing magnetic toner particles having an aimed particle size.

[0121] An important point of the present invention exists in that the black magnetic composite particles comprising the magnetic iron oxide particles which have an average particle size of 0.055 to 0.95 µm and may be coated with at least one compound selected from the group consisting of hydroxides of aluminum, oxides of aluminum, hydroxides of silicon and oxides of silicon; the organosilicon compounds coated on at least a part of the surface of the magnetic iron oxide particle; the carbon black fine particles having an average particle size of 0.002 to 0.05 µm, which are adhered on the surface of the coating composed of the organosilicon compounds, in which the total amount of the carbon black fine particles adhered to the coating composed of the organosilicon compounds is 1 to 25 parts by weight based on 100 parts by weight of the magnetic iron oxide particles, can show not only excellent fluidity and blackness, but also an excellent dispersibility in a binder resin upon the production of magnetic toner due to less amount of carbon black fine particles desorbed or fallen-off from the surfaces of the particles.

[0122] The reason why the amount of the carbon black fine particles desorbed or fallen-off from the surfaces of the black magnetic composite particles according to the present invention, is small, is considered as follows. That is, the surfaces of the magnetic iron oxide particles and the organosilicon compounds are strongly bonded to each other, so that the carbon black fine particles bonded to the surfaces of the magnetic iron oxide particles through the organosilicon compounds can be prevented from being desorbed from the magnetic iron oxide particles.

[0123] In particular, in the case of the alkoxysilane compounds (1) and the fluoroalkylsilane compounds (3), metallosiloxane bonds (=Si-O-M wherein M represents a metal atom contained in the black iron oxide particles, such as Si, Al, Fe or the like) are formed between the surfaces of the magnetic iron oxide particles and alkoxy groups contained in the organosilicon compounds onto which the carbon black fine particles are adhered, thereby forming a stronger bond bit with the organosilicon compounds on which the carbon black fine particles are adhered, and the surfaces of the magnetic iron oxide particles.

[0124] The reason why the black magnetic composite particles according to the present invention can show an accellent dispersibility in a binder resin upon the production of magnetic ton r, is considered such that since only a

small amount of the carbon black fine particles are desorbed or fallen-off from the surfaces of the black magnetic composite particles, the black magnetic composite particles is fre from deterioration in dispersibility due to the desorbed or fallen-off carbon black fine particles, and further since the carbon black fine particles are adhered onto the surfaces of the black magnetic composite particles and, therefore, irregularities are formed on the surfaces of the black magnetic composite particles, the contact between the particles can be suppressed.

[0125] The reason why the black magnetic composite particles according to the present invention can show an excellent fluidity, is considered as follows. That is, the carbon black fine particles which are ordinarily agglomerated together due to fineness thereof, are allowed to be uniformly and densely adhered on the surfaces of the magnetic iron oxide particles and, therefore, can be dispersed nearly in the form of primary particles, so that many fine irregularities are formed on the surfaces of the magnetic iron oxide particles.

[0126] The reason why the black magnetic composite particles according to the present invention can show an excellent blackness, is considered such that since the carbon black fine particles are uniformly and densely adhered on the surfaces of the magnetic iron oxide particles, the color tone of the core particles is hidden behind the carbon black fine particles, so that an inherent color tone of carbon black can be exhibited.

[0127] Therefore, the black magnetic toner produced by using the above black magnetic composite particles, can show excellent fluidity and blackness.

[0128] The reason why the black magnetic toner according to the present invention can show an excellent fluidity, is considered as follows. That is, the black magnetic composite particles on which a large amount of the carbon black fine particles are uniformly adhered, are blended in the black magnetic toner, so that many fine irregularities are formed on the surface of the black magnetic toner.

[0129] The reason why the black magnetic toner according to the present invention can show an excellent blackness, is considered such that the black magnetic composite particles having an excellent blackness is blended in the black magnetic toner.

[0130] As described above, since the black magnetic composite particles according to the present invention, are excellent not only in fluidity and blackness, but also in dispersibility in a binder resin due to less amount of the carbon black fine particles desorbed or fallen-off from the surfaces thereof, the black magnetic composite particles according to the present invention, are suitable as black magnetic particles for black magnetic toner capable of attaining a high image quality and a high copying speed.

[0131] In addition, since the black magnetic composite particles according to the present invention, are excellent in dispersibility in a binder resin, the particles can show excellent handling property and workability and, therefore, are preferable from an industrial viewpoint.

[0132] Further, the black magnetic toner produced from the above black magnetic composite particles which are excellent in fluidity and blackness, can also show excellent fluidity and blackness. Accordingly, the black magnetic toner is suitable as black magnetic toner capable of attaining a high image quality and a high copying speed.

[0133] Furthermore, in the black magnetic toner according to the present invention, since the black magnetic composite particles contained therein are excellent in dispersibility, it is possible to expose the black magnetic composite particles to the surface of the black magnetic toner independently and separately. As a result, the black magnetic toner can be free from being deteriorated in electric resistance due to the existence of the carbon black fine particles. Accordingly, the black magnetic toner according to the present invention is suitable as a high-resistance or insulated magnetic toner.

EXAMPLES

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[0134] The present invention is described in more detail by Examples and Comparative Examples, but the Examples are only illustrative and, therefore, not intended to limit the scope of the present invention.

[0135] Various properties were measured by the following methods.

- (1) The average particle size, the average major axial diameter and average minor axial diameter of magnetite particles, maghemite particles, black magnetic composite particles and carbon black fine particles were respectively expressed by the average of values (measured in a predetermined direction) of about 350 particles which were sampled from a micrograph obtained by magnifying an original electron micrograph (× 20,000) by four times in each of the longitudinal and transverse directions.
- (2) The <u>asp_ct ratio</u> of the particles was expr_ssed by the ratio of averag _major axial diameter to av_rag _minor axial diameter thereof.
- (3) The geometrical standard d viation of particle sizes was expressed by values obtained by the following method. That is, the particle sizes (major axial diameters) were measured from the above magnified electron micrograph. The actual particle sizes (major axial diameters) and the number of the particles were calculated from the measured values. On a logarithmic normal probability paper, the particle sizes (major axial diameters) were plotted at regular

intervals on the abscissa-axis and the accumulative number (under integration $si \ v$) of particles belonging to each interval of the particle $siz \ s$ (major axial diamet rs) w re plotted by percentage on the ordinate-axis by a statistical technique.

[0136] The particle sizes (major axial diameters) corresponding to the number of particles of 50 % and 84.13 %, respectively, were read from the graph, and the geometrical standard deviation was calculated from the following formula:

Geometrical standard deviation =

(particle size (major axial diameters)

corresponding to 84.13 % under integration

sieve}/{particle size (major axial diameters)

(geometrical average diameter) corresponding

to 50 % under integration sieve)

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[0137] The closer to 1 the geometrical standard deviation value, the more excellent the particle size distribution.

- (4) The specific surface area was expressed by the value measured by a BET method.
- (5) The <u>amounts of Al and Si</u> which were present within black magnetic composite particles or on surfaces thereof, and the <u>amount of Si</u> contained in the organosilicon compounds, were measured by a fluorescent X-ray spectroscopy device 3063 (manufactured by Rigaku Denki Kogyo Co., Ltd.) according to JIS K0119 "General rule of fluorescent X-ray analysis".
- (6) The <u>amount of carbon</u> adhered on the black magnetic composite particles was measured by "Horiba Metal, Carbon and Sulfur Analyzer EMIA-2200 Model" (manufactured by Horiba Seisakusho Co., Ltd.).
- (7) The <u>fluidity</u> of magnetic iron oxide particles, black magnetic composite particles and magnetic toner was expressed by a fluidity index which was a sum of indices obtained by converting on the basis of the same reference measured values of an angle of repose, a degree of compaction (%), an angle of spatula and a degree of agglomeration as particle characteristics which were measured by a powder tester (tradename, produced by Hosokawa Micron Co., Ltd.). The closer to 100 the fluidity index, the more excellent the fluidity of the particles.
- (8) The <u>blackness</u> of magnetic iron oxide particles, black magnetic composite particles and magnetic toner was measured by the following method. That is, 0.5 g of sample particles and 1.5 cc of castor oil were intimately kneaded together by a Hoover's muller to form a paste. 4.5 g of clear lacquer was added to the obtained paste and was intimately kneaded to form a paint. The obtained paint was applied on a cast-coated paper by using a 6-mil applicator to produce a coating film piece (having a film thickness of about 30 μm). The thus obtained coating film piece was measured according to JIS Z 8729 by a multi-light source spectrographic colorimeter MSC-IS-2D (manufactured by Suga Testing Machines Manufacturing Co., Ltd.) to determine an L* value of colorimetric indices thereof. The blackness was expressed by the L* value measured.

Here, the L* value represents a lightness, and the smaller the L* value, the more excellent the blackness.

(9) The <u>desorption percentage</u> of carbon black fine particles adhered on the black magnetic composite particles was measured by the following method. The closer to zero the desorption percentage, the smaller the amount of carbon black fine particles desorbed from the surfaces of black magnetic composite particles.

That is, 3 g of the black magnetic composite particles and 40 ml of ethanol were placed in a 50-ml precipitation pipe and then was subjected to ultrasonic dispersion for 20 minutes. Thereafter, the obtained dispersion was allowed to stand for 120 minutes, and the carbon black fine particles desorbed were separated from the black magnetic composite particles on the basis of the difference in specific gravity between both the particles. Next, the black magnetic composite particles from which the desorbed carbon black fine particles were separated, were mixed again with 40 ml of ethanol, and the obtained mixtur was furth r subject d to ultrasonic dispersion for 20 minut s. Ther after, the obtained dispersion was allowed to stand for 120 minutes, thereby separating the black magnetic composited particles and the desorbed carbon black desorbed from each other. The thus obtained black magnetic composited particles are dried at 100°C for one hour, and then the carbon content thereof was measured by the "Horiba Metal, Carbon and Sulfur Analyzer EMIA-2200 Model" (manufactured by Horiba Seisakusho Co., Ltd.). The desorption percentage of the carbon black fine particles was calculated according to the following for-

mula:

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Desorption percentage of

carbon black fine particles

$$= \{ (W_a - W_e)/W_a \} \times 100$$

wherein W_a represents an amount of carbon black fine particles initially adhered on the black magnetic composite particles; and W_e represents an amount of carbon black fine particles still adhered on the black magnetic composite particles after desorption test.

- (10) The <u>dispersibility</u> in a binder resin of the black magnetic composite particles was evaluated by counting the number of undispersed agglomerated particles on a micrograph (x 200 times) obtained by photographing a sectional area of the obtained black magnetic toner particle using an optical microscope (BH-2, manufactured by Olympus Kogaku Kogyo Co., Ltd.), and classifying the results into the following five ranks. The 5th rank represents the most excellent dispersing condition.
- Rank 1: not less than 50 undispersed agglomerated particles per 0.25 mm² were recognized;
- Rank 2: 10 to 49 undispersed agglomerated particles per 0.25 mm² were recognized;
- Rank 3: 5 to 9 undispersed agglomerated particles per 0.25 mm² were recognized;
- Rank 4: 1 to 4 undispersed agglomerated particles per 0.25 mm² were recognized;
- Rank 5: No undispersed agglomerated particles were recognized.

(11) The <u>average particle size</u> of the black magnetic toner was measured by a laser diffraction-type particle size distribution-measuring apparatus (Model HELOSLA/KA, manufactured by Sympatec Corp.).

(12) The volume resistivity of the magnetic iron oxide particles, the black magnetic composite particles and the black magnetic toner was measured by the following method.

That is, first, 0.5 g of a sample particles or toner to be measured was weighted, and press-molded at 140 Kg/cm² using a KBr tablet machine (manufactured by Simazu Seisakusho Co., Ltd.), thereby forming a cylindrical test piece.

Next, the thus obtained cylindrical test piece was exposed to an atmosphere maintained at a temperature of 25°C and a relative humidity of 60 % for 12 hours. Thereafter, the cylindrical test piece was set between stainless steel electrodes, and a voltage of 15V was applied between the electrodes using a Wheatstone bridge (TYPE2768, manufactured by Yokogawa-Hokushin Denki Co., Ltd.) to measure a resistance value R (Ω) .

The cylindrical test piece was measured with respect to an upper surface area A (cm²) and a thickness t_0 (cm) thereof. The measured values were inserted into the following formula, thereby obtaining a volume resistivity X (Ω -cm).

$$X (\Omega \cdot cm) = R \times (A/t_0)$$

(13) The <u>magnetic properties</u> of the magnetic iron oxide particles and the black magnetic composite particles were measured using a vibration sample magnetometer "VSM-3S-15" (manufactured by Toei Kogyo Co., Ltd.) by applying an external magnetic field of 10 kOe thereto. Whereas, the <u>magnetic properties</u> of the black magnetic toner were measured by applying external magnetic fields of 1 kOe and 10 kOe thereto.

Example 1:

<Production of black magnetic composite particles>

[0138] 20 kg of spherical magnetite particles iz : 0.23 µm; geometrical standard deviation value: 1.42; BET specific surface area value: 9.2 m²/g; blackness (L* value): 20.6; fluidity index: 35; coercive force value: 61 Oe; saturation magnetization value in a magnetic field of 10 kOe: 84.9 emu/g; residual magnetization value in a magnetic field of 10 kOe: 7.8 emu/g), were deagglomerated in 150 liters of pur water using a stirrer, and further passed through a "TK pipelin homomix r" (tradename, manufactured by Tokushu Kika Kogyo Co., Ltd.) three times, thereby obtaining a sturry containing the spherical magnetite particles.

[0139] Successively, the obtained slurry containing the spherical magnetite particles was pass of through a transverse-type sand grinder (tradename "MIGHTY MILL MHG-1.5L", manufactured by Inoue Seisakusho Co., Ltd.) five times at an axis-rotating speed of 2,000 rpm, thereby obtaining a slurry in which the spherical magnetit particles were dispersed.

[0140] The particles in the obtained slurry which remained on a sieve of 325 meshes (mesh size: 44 µm) was 0 %. The slurry was filtered and washed with water, thereby obtaining a filter cake containing the spherical magnetite particles. After the obtained filter cake containing the spherical magnetite particles was dried at 120°C, 11.0 kg of the dried particles were then charged into an edge runner "MPUV-2 Model" (tradename, manufactured by Matsumoto Chuzo Tekkosho Co., Ltd.), and mixed and stirred at 30 Kg/cm for 30 minutes, thereby lightly deagglomerating the particles. [0141] 110 g of methyl triethoxysilane was mixed and diluted with 200 ml of ethanol to obtain a methyl triethoxysilane

[0141] 110 g of methyl triethoxysilane was mixed and diluted with 200 ml of ethanol to obtain a methyl triethoxysilane solution. The methyl triethoxysilane solution was added to the deagglomerated spherical magnetite particles under the operation of the edge runner. The spherical magnetite particles were continuously mixed and stirred at a linear load of 60 Kg/cm for 60 minutes.

[0142] Next, 990 g of carbon black fine particles shown in the electron micrograph (× 20,000) of Fig. 2 (particle shape: granular shape; average particle size: 0.022 μm; geometrical standard deviation value: 1.68; BET specific surface area value: 134 m²/g; and blackness (L* value): 16.6) were added to the spherical magnetite particles coated with methyl triethoxysilane for 10 minutes while operating the edge runner. Further, the mixed particles were continuously stirred at a linear load of 60 Kg/cm for 60 minutes to adhere the carbon black fine particles on the coating composed of methyl triethoxysilane, thereby obtaining black magnetic composite particles.

[0143] The obtained black magnetic composite particles were aged at 105°C for 60 minutes by using a drier to evaporate water, ethanol or the like which were remained on surfaces of the composite particles. As shown in the electron micrograph (× 20,000) of Fig. 3, the resultant black magnetic composite particles had an average particle size of 0.24 μm. In addition, the black magnetic composite particles showed a geometrical standard deviation value of 1.42, a BET specific surface area value of 10.2 m²/g, a fluidity index of 46 and a blackness (L* value) of 18.5. The desorption percentage of the carbon black fine particles from the black magnetic composite particles was 7.5 %. As to the magnetic properties, the coercive force value of the black magnetic composite particles was 61 Oe; the saturation magnetization value in a magnetic field of 10 kOe was 77.3 emu/g; and the residual magnetization value in a magnetic field of 10 kOe was 7.1 emu/g. The coating amount of an organosilane compound produced from methyl triethoxysilane was 0.31 % by weight calculated as Si. Since no independent carbon black fine particles were observed on the electron micrograph of Fig. 3, it was determined that a whole amount of the carbon black fine particles were adhered on the coating composed of the organosilane compound produced from methyl triethoxysilane.

Example 2:

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35 < Production of black magnetic toner containing black magnetic composite particles>

[0144] 400 g of the black magnetic composite particles obtained in Example 1, 540 g of styrene-butyl acrylate-methyl methacrylate copolymer resin (molecular weight = 130,000, styrene/butyl acrylate/methyl methacrylate = 82.0/16.5/1.5), 60 g of polypropylene wax (molecular weight: 3,000) and 15 g of a charge-controlling agent were charged into a Henschel mixer, and mixed and stirred therein at 60°C for 15 minutes. The obtained mixed particles were melt-kneaded at 140°C using a continuous-type twin-screw kneader (T-1), and the obtained kneaded material was cooled, coarsely pulverized and finely pulverized in air. The obtained particles were subjected to classification, thereby producing a black magnetic toner.

[0145] The obtained black magnetic toner had an average particle size of 9.7 μm, a dispersibility of 5th rank, a fluidity index of 73, a blackness (L* value) of 18.3, a volume resistivity of 1.0 × 10¹⁴ Ω•cm, a coercive force value of 60 Oe, a saturation magnetization value (in a magnetic field of 10 kOe) of 32.6 emu/g, a residual magnetization value (in a magnetic field of 1 kOe) of 25.9 emu/g, and a residual magnetization value (in a magnetic field of 1 kOe) of 3.5 emu/g.

Example 3:

<Production of black magnetic composite particles>

[0146] 20 kg of spherical magnetite particles shown in the electron micrograph (× 20,000) of Fig. 1 (av rage particle size: 0.23 µm; geometrical standard d viation value: 1.42; BET specific surface ar a value: 9.2 m²/g; blackness (L* value): 20.6; fluidity ind x: 35; coerciv force value: 61 Oe; saturation magnetization valu in a magnetic field of 10 kO: 84.9 emu/g; residual magnetization valu in a magnetic fi ld of 10 kO: 7.8 emu/g), w r deagglomerated in 150 liters of pure water using a stirrer, and furth r passed through a "TK pipeline homomixer" (tradenam, manufactured

by Tokushu Kika Kogyo Co., Ltd.) three times, thereby obtaining a slurry containing the sph rical magn tite particles. [0147] Successiv ly, the obtain d slurry containing the spherical magnetite particles was passed through a transverse-type sand grind r (trad name "MIGHTY MILL MHG-1.5L", manufactured by Inoue S isakusho Co., Ltd.) five times at an axis-rotating speed of 2,000 rpm, thereby obtaining a slurry in which the spherical magnetit particles were dispersed.

[0148] The particles in the obtained slurry which remained on a sieve of 325 meshes (mesh size: 44 µm) was 0 %. The slurry was filtered and washed with water, thereby obtaining a filter cake containing the spherical magnetite particles. After the obtained filter cake containing the spherical magnetite particles was dried at 120°C, 11.0 kg of the dried particles were then charged into an edge runner "MPUV-2 Model" (tradename, manufactured by Matsumoto Chuzo Tekkosho Co., Ltd.), and mixed and stirred at 30 Kg/cm for 30 minutes, thereby lightly deagglomerating the particles. [0149] 110 g of methyl hydrogen polysiloxane (tradename: "TSF484", produced by TOSHIBA SILICONE CO., LTD.) were added to the deagglomerated spherical magnetite particles under the operation of the edge runner. The spherical magnetite particles were continuously mixed and stirred at a linear load of 60 Kg/cm for 60 minutes.

[0150] Next, 990 g of carbon black fine particles shown in the electron micrograph (× 20,000) of Fig. 2 (particle shape: granular shape; average particle size: 0.022 µm; geometrical standard deviation value: 1.68; BET specific surface area value: 134 m²/g; and blackness (L* value): 16.6) were added to the spherical magnetite particles coated with methyl hydrogen polysiloxane for 10 minutes while operating the edge runner. Further, the mixed particles were continuously stirred at a linear load of 60 Kg/cm for 60 minutes to adhere the carbon black fine particles on the coating composed of methyl hydrogen polysiloxane, thereby obtaining black magnetic composite particles.

[0151] The obtained black magnetic composite particles were dried at 105°C for 60 minutes by using a drier to evaporate water or the like which were remained on surfaces of the composite particles. The obtained black magnetic composite particles had an average particle size of 0.24 μm. In addition, the black magnetic composite particles had a geometrical standard deviation value of 1.42, a BET specific surface area value of 9.8 m²/g, a fluidity index of 48 and a blackness (L*value) of 18.2. The desorption percentage of the carbon black fine particles from the black magnetic composite particles was 6.5 %. As to the magnetic properties, the coercive force value of the black magnetic composite particles was 59 Oe; the saturation magnetization value in a magnetic field of 10 kOe was 7.0 emu/g. The coating amount of methyl hydrogen polysiloxane was 0.44 % by weight calculated as Si. Since no independent carbon black fine particles were observed on the electron micrograph, it was determined that a whole amount of the carbon black fine particles were adhered on the coating composed of methyl hydrogen polysiloxane.

Example 4:

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<Production of black magnetic toner containing black magnetic composite particles>

[0152] 400 g of the black magnetic composite particles obtained in Example 3, 540 g of styrene-butyl acrylate-methyl methacrylate copolymer resin (molecular weight = 130,000, styrene/butyl acrylate/methyl methacrylate = 82.0/16.5/1.5), 60 g of polypropylene wax (molecular weight: 3,000) and 15 g of a charge-controlling agent were charged into a Henschel mixer, and mixed and stirred therein at 60°C for 15 minutes. The obtained mixed particles were melt-kneaded at 140°C using a continuous-type twin-screw kneader (T-1), and the obtained kneaded material was cooled, coarsely pulverized and finely pulverized in air. The obtained particles were subjected to classification, thereby producing a black magnetic toner.

[0153] The obtained black magnetic toner had an average particle size of 9.7 μ m, a dispersibility of 5th rank, a fluidity index of 72, a blackness (L* value) of 18.1, a volume resistivity of 1.2 \times 10¹⁴ Ω •cm, a coercive force value of 59 Oe, a saturation magnetization value (in a magnetic field of 10 kOe) of 32.4 emu/g, a residual magnetization value (in a magnetic field of 1 kOe) of 4.1 emu/g, a saturation magnetization value (in a magnetic field of 1 kOe) of 25.7 emu/g, and a residual magnetization value (in a magnetic field of 1 kOe) of 3.4 emu/g.

Example 5:

<Production of black magnetic composite particles>

[0154] 20 kg of spherical magnetit particl s shown in the electron micrograph (× 20,000) of Fig. 1 (av rage particle siz : 0.23 μm; geometrical standard deviation value: 1.42; BET specific surface area value: 9.2 m²/g; blackness (L* valu): 20.6; fluidity index: 35; coercive forc value: 61 Oe; saturation magnetization valu in a magnetic fill of 10 kOe: 84.9 emu/g; residual magnetization value in a magn tic field of 10 kOe: 7.8 emu/g), were deagglomerated in 150 liters of pure water using a tirrir, and furthir passed through a "TK pipelin homomixer" (tradename, manufacturid by Tokushu Kika Kogyo Co., Ltd.) three times, thereby blaining a slurry containing the sphirical magnetite particles.

[0155] Successively, the obtained slurry containing the spherical magnetite particles was pass d through a transv rs -type sand grinder (tradename "MIGHTY MILL MHG-1.5L", manufactured by Inou Seisakusho Co., Ltd.) five times at an axis-rotating sp ed of 2,000 rpm, ther by obtaining a slurry in which the spherical magnetite particles were dispersed.

[0156] The particles in the obtained slurry which remained on a sieve of 325 meshes (mesh size: 44 µm) was 0 %. The slurry was filtered and washed with water, thereby obtaining a filter cake containing the spherical magnetite particles. After the obtained filter cake containing the spherical magnetite particles was dried at 120°C, 11.0 kg of the dried particles were then charged into an edge runner "MPUV-2 Model" (tradename, manufactured by Matsumoto Chuzo Tekkosho Co., Ltd.), and mixed and stirred at 30 Kg/cm for 30 minutes, thereby lightly deagglomerating the particles. [0157] 220 g of tridecafluorooctyl trimethoxysilane (tradename "TSL8257", produced by TOSHIBA SILICONE CO., LTD.) were added to the deagglomerated spherical magnetite particles under the operation of the edge runner. The spherical magnetite particles were continuously mixed and stirred at a linear load of 60 Kg/cm for 60 minutes.

[0158] Next, 990 g of carbon black fine particles shown in the electron micrograph (× 20,000) of Fig. 2 (particle shape: granular shape; average particle size: 0.022 μm; geometrical standard deviation value: 1.68; BET specific surface area value: 134 m²/g; and blackness (L* value): 16.6) were added to the spherical magnetite particles coated with tridecafluorooctyl trimethoxysilane for 10 minutes while operating the edge runner. Further, the mixed particles were continuously stirred at a linear load of 60 Kg/cm for 60 minutes to adhere the carbon black fine particles on the coating composed of tridecafluorooctyl trimethoxysilane, thereby obtaining black magnetic composite particles.

[0159] The obtained black magnetic composite particles were aged at 105°C for 60 minutes by using a drier to evaporate water or the like which were remained on surfaces of the composite particles. The obtained black magnetic composite particles had an average particle size of 0.24 µm. In addition, the black magnetic composite particles showed a geometrical standard deviation value of 1.42, a BET specific surface area value of 8.6 m²/g, a fluidity index of 48 and a blackness (L*value) of 18.4. The desorption percentage of the carbon black fine particles from the black magnetic composite particles was 6.8 %. As to the magnetic properties, the coercive force value of the black magnetic composite particles was 61 Oe; the saturation magnetization value in a magnetic field of 10 kOe was 76.8 emu/g; and the residual magnetization value in a magnetic field of 10 kOe was 6.9 emu/g. The coating amount of a fluorine-containing organosilane compound produced from tridecafluorooctyl trimethoxysilane was 0.13 % by weight calculated as Si. Since no independent carbon black fine particles were observed on the electron micrograph, it was determined that a whole amount of the carbon black fine particles were adhered on the coating composed of the fluorine-containing organosilane compound produced from tridecafluorooctyl trimethoxysilane.

Example 6:

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<Production of black magnetic toner containing black magnetic composite particles>

[0160] 400 g of the black magnetic composite particles obtained in Example 5, 540 g of styrene-butyl acrylate-methyl methacrylate copolymer resin (molecular weight = 130,000, styrene/butyl acrylate/methyl methacrylate = 82.0/16.5/1.5), 60 g of polypropylene wax (molecular weight: 3,000) and 15 g of a charge-controlling agent were charged into a Henschel mixer, and mixed and stirred therein at 60°C for 15 minutes. The obtained mixed particles were melt-kneaded at 140°C using a continuous-type twin-screw kneader (T-1), and the obtained kneaded material was cooled, coarsely pulverized and finely pulverized in air. The obtained particles were subjected to classification, thereby producing a black magnetic toner.

[0161] The obtained black magnetic toner had an average particle size of 10.1 μ m, a dispersibility of 5th rank, a fluidity index of 75, a blackness (L* value) of 18.5, a volume resistivity of 1.3 \times 10¹⁴ Ω •cm, a coercive force value of 58 Oe, a saturation magnetization value (in a magnetic field of 10 kOe) of 32.4 emu/g, a residual magnetization value (in a magnetic field of 10 kOe) of 4.2 emu/g, a saturation magnetization value (in a magnetic field of 1 kOe) of 25.7 emu/g, and a residual magnetization value (in a magnetic field of 1 kOe) of 3.4 emu/g.

Core particles 1 to 4:

[0162] Various magnetic iron oxide particles were prepared by known methods. The same procedure as defined in Example 1 was conducted by using the thus prepared particles, thereby obtaining deagglomerated magnetic iron oxide particles as core particles.

[0163] Various prop rties of the thus obtained magnetic iron oxide particles are shown in Table 1.

Core particles 5:

[0164] The same proc dure as defined in Example 1 was conducted by using 20 kg of the deagglomerated octahedral

magnetite particles (core particl s 1) and 150 liters of water, thereby obtaining a sturry containing the octah dral magnetite particles. The pH value of the obtained re-dispersed sturry containing the octah dral magnetite particles was adjusted to 4.0, and then the concentration of the sturry was adjusted to 98 g/liter by adding water thereto. After 150 liters of the sturry was in ated to 60°C, 2722 ml of a 1.0 mol/liter aluminum sulfate solution (equivalent to 1.0 % by weight (calculated as Al) based on the weight of the octahedral magnetite particles) was added to the sturry. After allowing the sturry to stand for 30 minutes, the pH value of the sturry was adjusted to 7.5 by adding an aqueous sodium hydroxide solution. Successively, 254 g of water glass #3 (equivalent to 0.5 % by weight (calculated as SiO₂) based on the weight of the octahedral magnetite particles) was added to the sturry. After the sturry was aged for 30 minutes, the pH value of the sturry was adjusted to 7.5 by adding acetic acid. After further allowing the sturry to stand for 30 minutes, the sturry was subjected to filtration, washing with water, drying and pulverization, thereby obtaining the octahedral magnetite particles coated with hydroxides of aluminum and oxides of silicon.

[0165] Main production conditions are shown in Table 2, and various properties of the obtained octahedral magnetite particles are shown in Table 3.

15 Core particles 6 to 8:

[0166] The same procedure as defined in the production of the core particles 5 above, was conducted except that kind of core particles, and kind and amount of additives used in the surface treatment were varied, thereby obtaining surface-treated magnetic iron oxide particles.

20 [0167] Main production conditions are shown in Table 2, and various properties of the obtained surface-treated magnetic iron oxide particles are shown in Table 3.

Examples 7 to 14 and Comparative Examples 1 to 5:

25 [0168] The same procedure as defined in Example 1 was conducted except that kind of particles to be treated, addition or non-addition of an alkoxysilane compound in the coating treatment with alkoxysilane compound, kind and amount of the alkoxysilane compound added, treating conditions of edge runner in the coating treatment, kind and amount of carbon black fine particles adhered, and treating conditions of edge runner used in the adhering process of the carbon black fine particles, were varied, thereby obtaining black magnetic composite particles. The black magnetic composite particles obtained in Examples 7 to 14 were observed by an electron microscope. As a result, almost no independent carbon black fine particles were recognized. Therefore, it was confirmed that a substantially whole amount of the carbon black fine particles were adhered on the coating composed of organosilane compound produced from the alkoxysilane compound.

[0169] Various properties of the carbon black fine particles A to C are shown in Table 4.

[0170] Main production conditions are shown in Table 5, and various properties of the obtained black magnetic composite particles are shown in Table 6.

[0171] Meanwhile, in Comparative Example 1, the spherical magnetite particles uncoated with the alkoxysilane compound and the carbon black fine particles were mixed and stirred together by an edge runner in the same manner as described above, thereby obtaining treated particles. An electron micrograph (× 20,000) of the thus treated particles is shown in Fig. 4. As shown in Fig. 4, it was confirmed that the carbon black fine particles were not adhered on the surfaces of the spherical magnetite particles, and both the particles were present independently.

Examples 15 to 22 and Comparative Examples 6 to 14:

45 < Production of black magnetic toner>

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[0172] The same procedure as defined in Example 2 was conducted by using the black magnetic composite particles obtained in Examples 7 to 14, the magnetic iron oxide particles as core particles 1 to 4, the mixed particles composed of the magnetic iron oxide particles and the carbon black fine particles used in Comparative Example 1 and the black magnetic composite particles obtained in Comparative Examples 2 to 5, thereby obtaining black magnetic toners.

[0173] Main production conditions and various properties of the obtained black magnetic toners are shown in Tables 7 and 8.

Examples 23 to 46 and Comparative Examples 15 to 23:

[0174] The same procedure as defined in Example 3 was conducted exc pt that kind of particles to be treated, addition or non-addition of a polysiloxane or modified polysiloxane, kind and amount of the polysiloxane or modified polysiloxane, treating conditions of edg runn r in the coating treatment, kind and amount of carbon black fine particles

adhered, and tr ating conditions of edg runner us d in the adhering process of the carbon black fin particles, w re varied, thereby obtaining black magnetic composite particles. The black magnetic composite particles obtained in Examples 23 to 46 were observed by an I ctron microscop. As a result, almost no independent carbon black fine particles were recognized. The refore, it was confirmed that a substantially whole amount of the carbon black fine particles were adhered on the coating composed of polysiloxane or modified polysiloxane.

[0175] Main production conditions are shown in Tables 9 to 11, and various properties of the obtained black magnetic composite particles are shown in Tables 12 to 14.

Examples 47 to 70 and Comparative Examples 24 to 32:

<Production of black magnetic toner>

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[0176] The same procedure as defined in Example 4 was conducted by using the black magnetic composite particles obtained in Examples 47 to 70, and the black magnetic composite particles obtained in Comparative Examples 15 to 23, thereby obtaining black magnetic toners.

[0177] Main production conditions and various properties of the obtained black magnetic toners are shown in Tables 15 to 17.

Examples 71 to 78 and Comparative Examples 33 to 35:

[0178] The same procedure as defined in Example 5 was conducted except that kind of particles to be treated, addition or non-addition of a fluoroalkyl organosilane compound, kind and amount of the fluoroalkyl organosilane compound added, treating conditions of edge runner in the coating treatment, kind and amount of carbon black fine particles adhered, and treating conditions of edge runner used in the adhering process of the carbon black fine particles, were varied, thereby obtaining black magnetic composite particles. The black magnetic composite particles obtained in Examples 71 to 78 were observed by an electron microscope. As a result, almost no independent carbon black fine particles were recognized. Therefore, it was confirmed that a substantially whole amount of the carbon black fine particles were adhered on the coating composed of a fluorine-containing organosilane compound produced from the fluoroalkyl organosilane compound.

30 [0179] Main production conditions are shown in Table 18, and various properties of the obtained black magnetic composite particles are shown in Table 19.

Examples 79 to 86 and Comparative Examples 36 to 38:

35 < Production of black magnetic toner>

[0180] The same procedure as defined in Example 6 was conducted by using the black magnetic composite particles obtained in Examples 71 to 78, and the black magnetic composite particles obtained in Comparative Examples 33 to 35, thereby obtaining black magnetic toners.

40 [0181] Main production conditions and various properties of the obtained black magnetic toners are shown in Table 20.

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Table 1

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	Properties of magnetic iron oxide particles				
Core	Kind	Particle	Average	Aspect	Geomet-
particles		shape	particle	ratio	rical
			size	(-)	standard
			(µm)	:	deviation
					(-)
Core	Magnetite	Octa-	0.28	•	1.53
particles	particles	hedral			
1					
Core	Magnetite	Spherical	0.23		1.35
particles	particles				
2					
Core	Magnetite	Acicular	0.40	8.1:1	1.53
particles	particles				
3					
Core	Maghemite	Spherical	0.20	_	1.42
particles	particles				
4			<u> </u>		

Table 1 (continued)

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	Proper	operties of magnetic iron oxide particles					
	BET	Magr	etic prop	perties			
Core	specific	Coer-	Satura-	Resid-	Fluid-	Black-	
particles	surface	cive	tion	ual	ity	ness	
	area	force	magnet-	magnet-	index	(L*	
	(m ² /g)	(0e)	ization	ization	(-)	value)	
			(10k0e)	(10k0e)		(-)	
			(emu/g)	(emu/g)			
Core	4.6	101	86.8	12.2	34	20.3	
particles							
11							
Core	11.8	63	85.1	7.7	38	20.1	
particles							
2							
Core	18.8	343	86.3	29.3	32	23.8	
particles							
3					_		
Core	7.2	54	78.8	8.7	38	31.5	
particles					·		
4							

Kind of

core

particles

Core

particles 1

Core

particles 2

Core

particles 3

Core

particles 4

Core

particles

Core

particles 5

Core

particles 6

Core

particles 7

Core

particles 8

Table 2

Kind

Aluminum

sulfate Water glass

#3

Sodium

aluminate Colloidal

silica

Aluminum

acetate

Water glass

#3

Surface-treating process

Additives

Calcu-

lated as

Al

 SiO_2

Al

 SiO_2

Al

SiO2

Amount

(wt. %)

1.0

0.5

2.0

1.0

5.0

1.0

5	

10

15

20

25

Table 2 (continued)

30

35

40

45

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55

	Surface-treating process			
Core		Coating materia	al	
particles	Kind	Calculated	Amount	
		as	(wt. %)	
Core particles 5	A	Al	0.98	
	S	SiO ₂	0.49	
Core particles 6	A	Al	1.92	
	s	SiO ₂	0.96	
Core particles 7	A	Al	4.75	
Core particles 8	S	SiO ₂	0.98	

Note; A: Hydroxide of aluminum

S: Oxide of silicon

Table 3

_						
10	Core particles	Properties of surface-treated magnetic iron oxide particles				
		Average	Aspect	Geometrical	BET	
		particle	ratio	standard	specific	
15		size	(-)	deviation	surface	
		(mu)		(-)	area	
				•	(m ² /g)	
20	Core	0.29	-	1.51	9.8	
	particles					
25	5					
	Core	0.24	-	1.35	13.6	
	particles					
30	6					
	Core	0.40	8.1:1	1.52	25.4	
35	particles		!			
	7					
40	Core	0.20	-	1.42	7.5	
	particles			•		
	8					

Table 3 (continued)

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15	

	Properties of surface-treated magnetic iron oxide particles				
Core	Magne	etic prope			
particles	Coercive	Satura-	Residual	Fluidity	Blackness
	force	tion	magnet-	index	(L*
	(0e)	magnet-	ization	(-)	value)
		ization	(10k0e)		(-)
		(10k0e)	(emu/g)		
		(emu/g)		•	
Core	103	86.3	12.1	32	21.4
particles					
5					
Core	62	84.8	7.6	37	20.8
particles					
- 6					
Core	336	86.0	19.8	32	24.6
particles					
7			<u> </u>		
Core	53	78.6	8.6	37	31.6
particles					
8				ł 	

Particle

shape

Granular

Granular

Granular

Table 4

Properties of carbon black fine particles

Average

particle

size

(µm)

0.022

0.015

0.030

Geometrical

standard

deviation

(-)

1.78

1.56

2.06

5

10

Kind of carbon

black fine

particles

Carbon black A

Carbon black B

Carbon black C

15

20

Table 4 (continued)

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Properties of carbon black fine Kind of carbon particles black fine BET specific Blackness surface area particles (L* value) (-) (m^2/g) Carbon black A 133.5 14.6 Carbon black B 265.3 15.2 Carbon black C 84.6 17.0

Table 5

,				
5				
			Production of black	
	Examples	Kind of	composite partic	
	and	particles	Coating step with alk	
10	Comparative	to be	or silicon compo	ound
	Examples	treated	Alkoxysilane comp	ound
				Amount
			Kind	added
15				(part by
15				weight)
	Example 7	Core	Dimethyl	1.0
		particles 1	dimethoxysilane	
	Example 8	Core	Phenyl	0.5
20		particles 2	triethoxysilane	
	Example 9	Core	Isobutyl	5.0
		particles 3	trimethoxysilane	
	Example 10	Core	Methyl	1.5
25		particles 4	triethoxysilane	
20	Example 11	Core	Dimethyl	0.2
	·	particles 5	dimethoxysilane	
	Example 12	Core	Phenyl	1.5
		particles 6	triethoxysilane	1
30	Example 13	Core	Isobutyl	1.0
		particles 7	trimethoxysilane	
	Example 14	Core	Methyl	2.0
!		particles 8	triethoxysilane	
35	Comparative	Core	=	_
	Example 1	particles		
		used in		
		Example 1		
40	Comparative	Core	Methyl	1.0
40	Example 2	particles 1	triethoxysilane	
	Comparative	Core	Dimethyl	0.5
	Example 3	particles 1	dimethoxysilane	
	Comparative	Core	Methyl	0.005
45	Example 4	particles 1	triethoxysilane	
	Comparative	Core	γ-aminopropyl	1.0
	Example 5	particles 1	triethoxysilane	

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Table 5 (continued)

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Examples	Production of black magnetic composite		
and	particles Coating step with alkoxysilane or silicon		
Comparative	coating step	compound	ne or silicon
Examples	Edge runne	r treatment	Coating amount
	Linear load	Time	(calculated as
	(Kg/cm)	(min)	Si)
			(wt. 왐)
Example 7	45	15	0.22
Example 8	75	20	0.06
Example 9	30	60 ·	0.73
Example 10	60	30	0.24
Example 11	60	20	0.05
Example 12	30	60	0.18
Example 13	45	30	0.16
Example 14	60	30	0.32
Comparative	-	-	-
. Example 1			
Comparative	30	60	0.21
Example 2			
Comparative	60	30	0.11
Example 3			
Comparative	60	30	7.9×10 ⁻⁴
Example 4			
Comparative	60	60	0.126
Example 5			

Table 5 (continued)

Production of black magnetic composite

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Examples particles and Adhering step of carbon black fine particles Comparative Carbon black fine particles Examples Kind Amount added (part by weight) Example 7 В 6.0 Example 8 В 12.0 Example 9 С 16.0 Example 10 Ä 25.0 Example 11 В 20.0 Example 12 В 15.0 Example 13 C 10.0 Example 14 A 20.0 Comparative Carbon black fine 10.0 Example 1 particles used in Example 1 Comparative Example 2 Comparative A 0.01 Example 3 Comparative В 5.0 Example 4 Comparative C 7.0 Example 5

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Table 5 (continued)

Production of black magnetic composite

Examples	particles		
and	Adhering step o	fine particles	
Comparative	Edge runne:	r treatment	Amount adhered
Examples	Linear load	Time	(calculated
	(Kg/cm)	(min)	as C)
			(wt. %)
Example 7	30	60	5.66
Example 8	30	90 .	10.73
Example 9	45	45	13.70
Example 10	60	60	22.65
Example 11	30	45	16.63
Example 12	60	60	12.99
Example 13	60	30	9.09
Example 14	75	30	17.09
Comparative	60	30	9.06
Example 1			
Comparative	-	-	-
Example 2			
Comparative	30 -	60	0.01
Example 3			
Comparative	60	45	4.75
Example 4			
Comparative	60	30	2.88
Example 5			

Table 6

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	Properties of black magnetic composite			
Examples	particles			
and	Average	Aspect	Geometrical	BET
Comparative	particle	ratio	standard	specific
Examples	size	(-)	deviation	surface
	(mrd)		(-)	area
				(m ² /g)
Example 7	0.28	-	1.52	5.0
Example 8	0.24	-	1.34	13.6
Example 9	0.41	8.1:1	1.51	23.8
Example 10	0.23	-	1.43	15.3
Example 11	0.30	_	1.47	14.4
Example 12	0.24	-	1.34	16.1
Example 13	0.40	8.0:1	1.50	24.8
Example 14	0.23	-	1.42	13.8
Comparative	0.29	-	1.53	11.9
Example 1			·	
Comparative	0.29	-	1.52	10.6
Example 2				
Comparative	0.28	-	1.52	5.6
Example 3				
Comparative	0.28	<u>-</u>	1.52	17.6
Example 4				
Comparative	0.29	-	1.52	11.2
Example 5				

Table 6 (continued)

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	Properties of black magnetic composite		
Examples	particles		
and	<u></u>	agnetic properti	
Comparative	Coercive	Saturation	Residual
Examples	force	magnetization	magnetization
	(0e)	(10 kOe)	(10 kOe)
		(emu/g)	(emu/g)
Example 7	108	81.1	11.4
Example 8	65	71.8	6.5
Example 9	336	73.8	25.8
Example 10	58	63.6	6.4
Example 11	106	72.8	10.2
Example 12	68	74.1	6.7
Example 13	331	77.8	27.2
Example 14	57	65.6	7.2
Comparative	103	79.3	10.3
Example 1			
Comparative	103	83.6	10.8
Example 2			
Comparative	104	86.7	11.3
Example 3			
Comparative	100	83.8	10.9
Example 4			
Comparative	102	84.6	10.6
Example 5		-	

5

Table 6 (continued)

Examples and Comparative index (L* value) (E* value) (R*) Example 7		Properties of black magnetic composite		
Comparative Examples	Examples	particles		
Examples (-) (-) percentage (%) Example 7 49 17.0 8.6 Example 8 45 16.4 8.2 Example 9 46 17.8 6.4 Example 10 54 17.5 5.2 Example 11 52 15.9 3.1 Example 12 47 16.2 3.6 Example 13 48 17.5 2.1 Example 14 51 17.9 3.8 Comparative 42 20.0 78.6 Example 1 Comparative 40 20.9 - Example 3 Comparative 40 20.9 - Example 3 Comparative 40 20.1 26.5 Example 4 Comparative 40 20.1 26.5	1		Blackness	Carbon black
Example 7 49 17.0 8.6 Example 8 45 16.4 8.2 Example 9 46 17.8 6.4 Example 10 54 17.5 5.2 Example 11 52 15.9 3.1 Example 12 47 16.2 3.6 Example 13 48 17.5 2.1 Example 14 51 17.9 3.8 Comparative 42 20.0 78.6 Example 1 Comparative 40 20.9 - Example 3 Comparative 40 20.9 Example 3 Comparative 40 20.1 26.5 Example 4 Comparative 41 20.6 41.6	, -		(L* value)	desorption
Example 7 49 17.0 8.6 Example 8 45 16.4 8.2 Example 9 46 17.8 6.4 Example 10 54 17.5 5.2 Example 11 52 15.9 3.1 Example 12 47 16.2 3.6 Example 13 48 17.5 2.1 Example 14 51 17.9 3.8 Comparative 42 20.0 78.6 Example 1 Comparative 40 20.9 - Example 3 Comparative 38 21.4 31.2 Example 4 Comparative 40 20.1 26.5 Example 4 Comparative 41 20.6 41.6	Examples	(-)	(-)	percentage
Example 8 45 16.4 8.2 Example 9 46 17.8 6.4 Example 10 54 17.5 5.2 Example 11 52 15.9 3.1 Example 12 47 16.2 3.6 Example 13 48 17.5 2.1 Example 14 51 17.9 3.8 Comparative 42 20.0 78.6 Example 1 Comparative 40 20.9 - Example 2 Comparative 38 21.4 31.2 Example 3 Comparative 40 20.1 26.5 Example 4 Comparative 41 20.6 41.6				(%)
Example 9 46 17.8 6.4 Example 10 54 17.5 5.2 Example 11 52 15.9 3.1 Example 12 47 16.2 3.6 Example 13 48 17.5 2.1 Example 14 51 17.9 3.8 Comparative 42 20.0 78.6 Example 1 Comparative 40 20.9 - Example 2 Comparative 38 21.4 31.2 Example 3 Comparative 40 20.1 26.5 Example 4 Comparative 41 20.6 41.6		49	17.0	8.6
Example 10 54 17.5 5.2 Example 11 52 15.9 3.1 Example 12 47 16.2 3.6 Example 13 48 17.5 2.1 Example 14 51 17.9 3.8 Comparative 42 20.0 78.6 Example 1 Comparative 40 20.9 - Example 2 Comparative 38 21.4 31.2 Example 3 Comparative 40 20.1 26.5 Example 4 Comparative 41 20.6 41.6				8.2
Example 11 52 15.9 3.1 Example 12 47 16.2 3.6 Example 13 48 17.5 2.1 Example 14 51 17.9 3.8 Comparative 42 20.0 78.6 Example 1 20.9 - Example 2 Comparative 38 21.4 31.2 Example 3 Comparative 40 20.1 26.5 Example 4 Comparative 41 20.6 41.6			17.8	6.4
Example 12 47 16.2 3.6 Example 13 48 17.5 2.1 Example 14 51 17.9 3.8 Comparative 42 20.0 78.6 Example 1 Comparative 40 20.9 - Example 2 Comparative 38 21.4 31.2 Example 3 Comparative 40 20.1 26.5 Example 4 Comparative 41 20.6 41.6				5.2
Example 13 48 17.5 2.1 Example 14 51 17.9 3.8 Comparative 42 20.0 78.6 Example 1 20.9 - Example 2 20.0 20.9 Comparative 38 21.4 31.2 Example 3 20.1 26.5 Example 4 20.6 41.6				3.1
Example 14 51 17.9 3.8 Comparative 42 20.0 78.6 Example 1 20.9 - Example 2 21.4 31.2 Example 3 20.1 26.5 Example 4 20.6 41.6	Example 12	47		3.6
Comparative 42 20.0 78.6 Example 1 20.9 - Comparative 40 20.9 - Example 2 21.4 31.2 Example 3 21.4 31.2 Comparative 40 20.1 26.5 Example 4 20.6 41.6	_		17.5	2.1
Example 1	Example 14	51	17.9	3.8
Comparative 40 20.9 -	Comparative	42	20.0	78.6
Example 2 Comparative 38 21.4 31.2 Example 3 Comparative 40 20.1 26.5 Example 4 Comparative 41 20.6 41.6	Example 1			
Comparative 38 21.4 31.2 Example 3 20.1 26.5 Example 4 20.6 41.6	Comparative	40	20.9	-
Example 3 Comparative 40 20.1 26.5 Example 4 Comparative 41 20.6 41.6	Example 2			
Comparative 40 20.1 26.5 Example 4 20.6 41.6	Comparative	38	21.4	31.2
Example 4 Comparative 41 20.6 41.6	Example 3			
Comparative 41 20.6 41.6	Comparative	40	20.1	26.5
1 2 1	Example 4			
From 1 o 5	Comparative	41	20.6	41.6
Example 2	Example 5			

Table 7

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Production of black magnetic toner Examples Black magnetic Resin composite particles Kind Amount Kind Amount blended blended (part by (part by weight) weight) Styrene-acryl Example 15 Example 7 45 55 copolymer resin Example 16 Example 8 45 Styrene-acryl 55 copolymer resin Example 17 Example 9 40 Styrene-acryl 60 copolymer resin Example 18 Example 10 50 Styrene-acryl 50 copolymer resin Example 19 Example 11 45 Styrene-acryl 55 copolymer resin Example 20 Example 12 40 Styrene-acryl 60 copolymer resin Example 21 Example 13 50 Styrene-acryl 50 copolymer resin Example 22 Example 14 50 Styrene-acryl 50 copolymer resin

Table 7 (continued)

Volume

resistivity

 $(\Omega \cdot cm)$

 9.8×10^{13}

1.6x10¹⁴

 7.3×10^{13}

 8.6×10^{13}

 6.8×10^{13}

 2.6×10^{14}

1.8x10¹⁴

1.1x10¹⁴

5					
		Properties of black magnetic toner			
	Examples	Average	Dispers-	Fluidity	Volum
10		particle	ibility	index	resisti
		size	(-)	(` -)	(Ω•cı
15		(µm)			
	Example 15	9.6	4	74	9.8x10
20	Example 16	10.1	5	82	1.6x10
	Example 17	11.2	4	72	7.3x10
25	Example 18	10.6	5	78	8.6x10
	Example 19	9.2	5	78	6.8x10
30	Example 20	9.8	5	86	2.6x10
	Example 21	8.9	5	73	1.8x10
35	Example 22	11.0	5	82	1.1x1(

36

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Table 7 (continued)

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	Propertie	s of black magne	etic toner
Examples	Magnetic properties		
	Coercive	Saturation m	agnetization
	force	(10 kOe) ·	(1 k0e)
	(Oe)	(emu/g)	(emu/g)
Example 15	96	36.8	27.6
Example 16	61	33.6	26.1
Example 17	311	31.4	23.4
Example 18	56	32.3	22.7
Example 19	103	32.3	23.6
Example 20	66	29.6	22.2
Example 21	320	39.1	29.7
Example 22	53	31.9	23.9

Table 7 (continued)

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	Properties of black magnetic toner		
Examples	Magnetic	properties	Blackness
	Residual ma	gnetization	(L* value)
	(10 kOe)	(1 kOe) .	(-)
	(emu/g)	(emu/g)	
Example 15	5.9	4.4	18.7
Example 16	4.0	2.9	18.1
Example 17	11.3	8.5	19.6
Example 18	4.5	3.2	19.2
- Example 19	5.2	4.0	17.4
Example 20	3.6	2.7	18.1
Example 21	14.1	9.9	18.9
Example 22	4.5	3.1	19.5

Table 8

_	-				
5		Production of black magnetic toner			toner
	Comparative	Black magnetic		Res	in
	Examples	partic	les		
		Kind	Amount	Kind	Amount
10			blended		blended
			(part by		(part by
			weight)		weight)
:	Comparative	Core	45	Styrene-	55
15	Example 6	particles		acryl	
		1		copolymer	
				resin	
	Comparative	Core	45	Styrene-	55
20	Example 7	particles		acryl	
20		2		copolymer	
	<u> </u>			resin	
	Comparative	Core	45	Styrene-	55
	Example 8	particles		acryl	
25		3		copolymer	
			4.5	resin	
	Comparative	Core	45	Styrene-	55
	Example 9	particles		acryl	
30		4		copolymer resin	
	Composition	Composition	45		55
	Comparative Example 10	Comparative Example 1	43	Styrene- acryl	23
	. Example 10	Example 1		copolymer	
				resin	
35	Comparative	Comparative	45	Styrene-	55
	Example 11	Example 2	=3	acryl	33
	manpic ii	Drampic 2		copolymer	
				resin	
40	Comparative	Comparative	45	Styrene-	55
	Example 12	Example 3		acryl	
				copolymer	
				resin	
45	Comparative	Comparative	45	Styrene-	55
	Example 13	Example 4		acryl	
	_	-		copolymer	
				resin	
50	Comparative	Comparative	45	Styrene-	55
50	Example 14	Example 5		acryl	
	_	_		copolymer	
				resin	

Table 8 (continued)

	Properties of black magnetic toner			
Comparative	Average	Dispers-	Fluidity	Volume
Examples	particle	ibility	index	resistivity
	size	(-)	(-)	(Ω•cm)
	(µm)			
Comparative	10.0	3	60	6.3x10 ¹²
Example 6				
Comparative	10.1	3	65	5.4x10 ¹²
Example 7				
Comparative	9.8	3	58	9.1x10 ¹¹
Example 8				
Comparative	10.3	3	63	3.2×10 ¹²
Example 9				
Comparative	11.0	2	55	3.6×10 ¹¹
Example 10				
Comparative	10.6	2	58	1.6x10 ¹²
Example 11				
Comparative	10.8	3	61	1.6x10 ¹²
Example 12				
Comparative	10.4	2	58	2.6x10 ¹²
Example 13		į		
Comparative	10.4	2	57	2.6x10 ¹²
Example 14				

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Table 8 (continued)

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Properties of black magnetic toner Comparative Magnetic properties Examples Coercive Saturation magnetization force (10 kOe) (1 kOe) (0e) (emu/g) (emu/g) Comparative 104 39.6 30.0 Example 6 Comparative 61 38.8 29.1 Example 7 Comparative 338 37.6 28.8 Example 8 Comparative 51 34.6 26.0 Example 9 Comparative 99 35.6 26.3 Example 10 Comparative 100 39.1 27.1 Example 11 Comparative 103 38.1 29.3 Example 12 Comparative 102 36.7 28.6 Example 13 Comparative 102 36.3 27.8 Example 14

Table 8 (continued)

	Properties of black magnetic toner		
Comparative	Magnetic	properties	Blackness
Examples	Residual ma	agnetization	(L* value)
	(10 kOe)	(1 k0e)	(-)
	(emu/g)	(emu/g)	
Comparative	5.5	4.2 .	22.3
Example 6		,	
Comparative	3.5	2.6	22.1
Example 7			
Comparative	12.8	9.7	26.0
Example 8			
Comparative	3.8	3.0	34.8
Example 9			
Comparative	5.0	3.7	22.6
Example 10			
Comparative	5.9	4.6	23.3
Example 11			
Comparative	5.3	4.1	23.5
Example 12			
Comparative	5.1	3.9	22.3
Example 13			
Comparative	4.9	3.8	22.1
Example 14			

Table 9

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Examples	Kind of	Production of black m	magnetic
and	particles	composite partic	les
Comparative	to be	Coating step with poly	
Examples	treated	Polysiloxane	
		Kind	Amount
			added
			(part by weight)
Example 23	Core	TSF484	1.0
		100 10 1	2.0
	particles 1		
Example 24	Core	TSF484	5.0
!	particles 2		
Example 25	Core	KF99	2.0
	particles 3		
Example 26	Core	L-9000	1.0
	particles 4		·
Example 27	Core	TSF451	1.5
	particles 5		
Example 28	Core	TSF484	3.5
	particles 6		
Example 29	Core	KF99	1.0
	particles 7		
Example 30	Core	L-9000	2.0
	particles 8		
Comparative	Core	TSF484	1.0
Example 15	particles 1		
Comparative	Core	TSF484	0.5
Example 16	particles 1		
Comparative	Core	TSF484	0.005
Example 17	particles 1		

Table 9 (continued)

	Production of black magnetic composite				
Examples		particles			
and	Coating	step with polys	siloxane		
Comparative	Edge runne	r treatment	Coating amount		
Examples	Linear load	Time	(calculated as		
	(Kg/cm)	(min)	Si)		
			(wt. %)		
Example 23	60	30	0.44		
Example 24	45	25	2.18		
Example 25	30	30	0.87		
Example 26	60	45	0.44		
Example 27	45	60	0.62		
Example 28	60	30	1.50		
Example 29	75	25	0.43		
Example 30	60	20	0.87		
Comparative	60	30	0.44		
Example 15					
Comparative	60	30	0.21		
Example 16					
Comparative	60	30	2.2x10 ⁻³		
Example 17					

Table 9 (continued)

	Production of black magnetic composite		
Examples	part	icles	
and	Adhering step of carbon black fine particles		
Comparative	Carbon black fine particles		
Examples	Kind	Amount added	
		(part by weight)	
Example 23	Α	. 10.0	
Example 24	A	3.0	
Example 25	В	5.0	
Example 26	С	10.0	
Example 27	A	5.0	
Example 28	A	10.0	
Example 29	В	15.0	
Example 30	С	10.0	
Comparative	-	_	
Example 15			
Comparative	A	0.01	
Example 16			
Comparative	В	3.0	
Example 17			
	and Comparative Examples Example 23 Example 24 Example 25 Example 26 Example 27 Example 28 Example 29 Example 30 Comparative Example 15 Comparative Example 16 Comparative	Examples and Adhering step of carbo Comparative Carbon black Examples Kind Example 23 A Example 24 A Example 25 B Example 26 C Example 27 A Example 28 A Example 29 B Example 30 C Comparative Example 15 Comparative A Example 16 Comparative B	

Table 9 (continued)

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5	

	Production of black magnetic composite		
Examples		particles	
and	Adhering step of	of carbon black	fine particles
Comparative	Edge runne	r treatment	Amount adhered
Examples	Linear load	Time	(calculated
	(Kg/cm)	(min)	as C)
		•	(wt. %)
Example 23	60	30	9.05
Example 24	60	45	2.89
Example 25	60	30	4.76
Example 26	45	45	9.12
Example 27	45	30	4.72
Example 28	30	60	8.99
Example 29	60	30	12.89
Example 30	45	25	9.08
Comparative	-	_	-
Example 15			
Comparative	60	30	0.01
Example 16			
Comparative	60	30	2.91
Example 17			

Table 10

5				
3	Examples	Kind of	Production of black r composite partic	les
	and	particles	Coating step with mo	odified
10	Comparative	to be	polysiloxane	
,,	Examples	treated	Modified polysilo	
			Kind	Amount added
				(part by
15				weight)
	Example 31	Core	BYK-080	1.0
		particles 1		
20	Example 32	Core	BYK-080.	0.5
		particles 2		
	Example 33	Core	BYK-310	2.0
25		particles 3		
	Example 34	Core	BYK-322	5.0
		particles 4		
30	Example 35	Core	BYK-080	2.0
		particles 5		
	Example 36	Core	BYK-080	3.0
35		particles 6		
	Example 37	Core	BYK-310	1.5
		particles 7		
40	Example 38	Core	BYK-322	7.0
_		particles 8		
<u>-</u>	Comparative	Core	BYK-080	1.0
45	Example 18	particles 1		
	Comparative	Core	BYK-080	0.5
50	Example 19	particles 1		
	Comparative	Core	BYK-080	0.005
	Example 20	particles 1		

Table 10 (continued)

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	Production of black magnetic composite				
Examples		particles			
and	Coating ster	with modified	polysiloxane		
Comparative	Edge runne:	r treatment	Coating amount		
Examples	Linear load	Time	(calculated		
	(Kg/cm)	(min)	as Si)		
		•	(wt. %)		
Example 31	60	60	0.18		
Example 32	30	60	0.08		
Example 33	60	45	0.36		
Example 34	30	30	0.87		
Example 35	45	30	0.36		
Example 36	45	45	0.49		
Example 37	60	30	0.25		
Example 38	30	45	1.20		
Comparative	60	30	0.16		
Example 18					
Comparative	60	30	0.08		
Example 19					
Comparative	60	30	8.0x10 ⁻⁴		
Example 20					

Table 10 (continued)

5			
		Production of black	magnetic composite
•	Examples	part.	icles
10	and	Adhering step of carbo	n black fine particles
	Comparative	Carbon black	fine particles
	Examples	Kind	Amount added
15			(part by weight)
	Example 31	A	. 8.0
	Example 32	A	6.0
20	Example 33	В	6.5
	Example 34	С	11.5
	Example 35	A	7.5
25	Example 36	A	12.5
	Example 37	В	18.0
	.Example 38	С	15.0
30	Comparative	-	-
	Example 18		
	Comparative	A	0.01
35	Example 19		
	Comparative	В	5.0
	Example 20		
	Example 20		

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Table 10 (continued)

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]	Production of black magnetic composite			
Examples		particles		
and	Adhering step	of carbon black	fine particles	
Comparative	Edge runne	r treatment	Amount adhered	
Examples	Linear load	Time	(calculated	
Ì	(Kg/cm)	(min)	as C)	
		•	(wt. %)	
Example 31	60	30	7.43	
Example 32	30	25	5.68	
Example 33	30	30	6.10	
Example 34	60	20	10.24	
Example 35	45	45	6.98	
Example 36	60	30	11.10	
Example 37	30	25	15.16	
Example 38	45	40	13.10	
Comparative	-	-	_	
Example 18				
Comparative	60	30	0.01	
Example 19				
Comparative	60	30	4.75	
Example 20				

Table 11

_				
5			Production of black r	_
	Examples	Kind of	composite partic	
	and Comparative	particles to be	Coating step with te modified polysilo	
10	Examples	treated	Terminal-modified poly	
10		02 0000	Kind	Amount
				added
				(part by
15				weight)
,,	Example 39	Core	TSF4770	2.0
		particles 1		
		_	man4550	
20	Example 40	Core	TSF4770.	1.0
		particles 2		
	Example 41	Core	TSF4751	0.5
		particles 3		
25				
	Example 42	Core	TSF4751	3.0
		particles 4		
30	Example 43	Core	TSF4770	1.0
		particles 5		
	Example 44	Core	TSF4770	3.0
		mambialas 6		3,73
35		particles 6		
	Example 45	Core	TSF4751	0.5
		particles 7		
40	Example 46	Core	TSF4751	1.7
	_	particles 8		
		•		
	Comparative	Core	TSF4770	1.0
45	Example 21	particles 1		
	Comparative	Core	TSF4770	1.0
	Example 22	particles 1		
50	Comparative	Core	TSF4770	0.005
	Example 23	particles 1		

Table 11 (continued)

5				
	Examples	Production o	of black magneti	ic composite
10	and Comparative		ep with termina polysiloxane	l-modified
	Examples	Edge runner	treatment	Coating amount
	Examples	Linear load	Time	(calculated
15		(Kg/cm)	(min) .	as Si)
				(wt. %)
	Example 39	60	30	0.46
20	Example 40	30	40	0.21
	Example 41	60	30	0.12
	Example 42	30	45	0.71
25	Example 43	45	20	0.21
	Example 44	60	30	0.69
	Example 45	45	20	0.14
30	Example 46	30	30	0.37
	Comparative	60	30	0.26
35	Example 21			
	Comparative	60	30	0.25
	Example 22			
40	Comparative	60	30	1.2x10 ⁻³
40	Example 23			

Table 11 (continued)

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	Production of black magnetic composite			
Examples	part	icles		
and	Adhering step of carbo	on black fine particles		
Comparative	Carbon black	fine particles		
Examples	Kind	Amount added		
		(part by weight)		
Example 39	A	10.0		
Example 40	A	6.0		
Example 41	В	8.0		
Example 42	С	10.0		
Example 43	A	7.5		
Example 44	A	12.0		
Example 45	В	19.0		
.Example 46	С	13.0		
Comparative	_	_		
Example 21				
Comparative	A	0.01		
Example 22	i i			
Comparative	В	5.0		
Example 23				

Table 11 (continued)

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	Production of black magnetic composite			
Examples		particles		
and	Adhering step	of carbon black	fine particles	
Comparative	Edge runne:	r treatment	Amount adhered	
Examples	Linear load	Time	(calculated	
	(Kg/cm)	(min)	as C)	
		•	(wt. %)	
Example 39	60	30	9.13	
Example 40	30	45	5.57	
Example 41	45	60	7.42	
Example 42	60	45	9.10	
Example 43	30	30	6.98	
Example 44	45	25	10.70	
Example 45	60	45	15.15	
Example 46	30	30	11.43	
Comparative	-	_	-	
Example 21				
Comparative	60	30	0.01	
Example 22				
Comparative	60	30	4.73	
Example 23				

Table 12

5				-	
	Examples	Properties of black magnetic composite particles			
10	and Comparative Examples	Average particle size (µm)	Aspect ratio (-)	Geometrical standard deviation (-)	BET specific surface area (m²/g)
15	Example 23	0.28	-	1.52	6.1
13	Example 24	0.24	-	1.34	12.8
	Example 25	0.41	8.1:1	1.51	24.6
20	Example 26	0.23	-	1.42	14.6
	Example 27	0.29	_	1.50	14.3
	Example 28	0.23	-	1.34	15.1
25	Example 29	0.40	8.1:1	1.50	24.8
	Example 30	0.23	_	1.42	12.8
30	Comparative Example 15	0.29	-	1.53	11.5
	Comparative	0.28	-	1.53	7.1
	Example 16				,
35	Comparative	0.28	_	1.53	15.6
	Example 17				

Table 12 (continued)

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	Properties of black magnetic composite			
Examples	particles			
and	Ma	agnetic properti	.es	
Comparative	Coercive	Saturation	Residual	
Examples	force	magnetization	magnetization	
	(0e)	(10 k0e)	(10 k0e)	
		(emu/g) ·	(emu/g)	
Example 23	108	80.8	11.0	
Example 24	64	71.6	6.6	
Example 25	338	75.2	25.8	
Example 26	56	64.1	6.3	
Example 27	105	73.1	10.1	
Example 28	65	75.0	6.3	
Example 29	336	77.6	26.5	
Example 30	58	65.1	6.8	
Comparative	103	78.6	10.1	
Example 15				
Comparative	104	83.6	10.9	
Example 16				
Comparative	103	83.2	10.6	
Example 17		·		

Table 12 (continued)

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	Properties of black magnetic composite			
Examples	particles			
and	Fluidity	Blackness	Carbon black	
Comparative	index	(L* value)	desorption	
Examples	(-)	(-)	percentage	
			(%)	
Example 23	51	17.0	7.2	
Example 24	48	16.5	8.6	
Example 25	46	17.2	8.8	
Example 26	53	17.4	6.2	
Example 27	50	15.3	4.6	
Example 28	48	16.0	3.6	
Example 29	49	17.6	1.8	
Example 30	51	17.6	3.2	
Comparative	39	20.6	60.5	
Example 15				
Comparative	39	21.0	28.3	
Example 16				
Comparative	40	20.8	43.8	
Example 17				

Table 13

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	Properties of black magnetic composite			
Examples		pa	rticles	
and	Average	Aspect	Geometrical	BET
Comparative	particle	ratio	standard	specific
Examples	size	(-)	deviation	surface
	(hw)		(-)	area
			•	(m^2/g)
Example 31	0.28	-	1.52	6.8
Example 32	0.23		1.34	11.9
Example 33	0.40	8.1:1	1.52	24.9
Example 34	0.23	-	1.42	13.8
Example 35	0.29	-	1.51	15.1
Example 36	0.23	-	1.34	14.6
Example 37	0.41	8.1:1	1.50	25.6
Example 38	0.23	-	1.42	11.8
Comparative	0.28	-	1.53	11.3
Example 18				
Comparative	0.28	-	1.52	10.6
Example 19				
Comparative	0.28	-	1.53	16.3
Example 20				
	<u> </u>	· · · · · · · · · · · · · · · · · · ·		

Table 13 (continued)

	Properties of black magnetic composite				
Examples	particles				
and	Magnetic properties				
Comparative	Coercive	Saturation	Residual		
Examples	force	magnetization	magnetization		
	(0e)	(10 kOe)	(10 k0e)		
		(emu/g) ·	(emu/g)		
Example 31	108	80.6	10.8		
Example 32	63	71.8	7.1		
Example 33	336	75.3	25.6		
Example 34	57	64.6	6.1		
Example 35	106	73.2	10.0		
Example 36	63	75.2	6.1		
Example 37	336	77.8	26.3		
Example 38	56	65.3	6.5		
Comparative	102	78.3	10.0		
Example 18					
Comparative	100	83.2	10.8		
Example 19		ĺ			
Comparative	102	81.6	10.1		
Example 20					

Table 13 (continued)

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	Properties of black magnetic composite				
Examples	particles				
and	Fluidity	Blackness	Carbon black		
Comparative	index	(L* value)	desorption		
Examples	(-)	(-)	percentage		
			(%)		
Example 31	52	17.1 ·	8.3		
Example 32	52	16.8	9.1		
Example 33	53	17.5	6.3		
Example 34	46	17.0	5.9		
Example 35	48	17.3	4.8		
Example 36	51	16.1	3.9		
Example 37	50	16.0	3.8		
Example 38	53	17.3	2.8		
Comparative	38	21.0	61.3		
Example 18					
Comparative	37	20.6	27.3		
Example 19					
Comparative	38	20.5	45.9		
Example 20					

Table 14

		5	.:	-1		
	Examples	Properties of black magnetic composite particles				
	and	Average	Aspect	Geometrical	BET	
	Comparative	particle	ratio	standard	specific	
)	Examples	size	(-)	deviation	surface	
		(mm)		(-)	area	
		0.00			(m ² /g)	
	Example 39	0.28	-	1.52	5.9	
•	Example 40	0.23	-	1.34	12.6	
	Example 41	0.41	8.0:1	1.52	25.6	
,	Example 42	0.23	-	1.43	13.9	
	Example 43	0.29	-	1.50	14.8	
	Example 44	0.23	-	1.34	16.3	
ī	Example 45	0.40	8.1:1	1.50	21.8	
	Example 46	0.23	-	1.43	13.6	
	Comparative	0.29	_	1.52	10.3	
)	Example 21					
	Comparative	0.29	-	1.51	10.6	
	Example 22					
ī	Comparative	0.29	-	1.52	10.1	
	Example 23					

Table 14 (continued)

5							
	_ ,	Properties of black magnetic composite					
	Examples and		particles				
			Magnetic properties				
10	Comparative	Coercive	Saturation	Residual			
	Examples	force	magnetization	magnetization			
		(0e)	(10 kOe)	(10 k0e)			
			(emu/g) ·	(emu/g)			
15	Example 39	107	81.4	11.6			
,3	Example 40	66	72.1	6.3			
	Example 41	341	74.1	25.4			
20	Example 42	59	63.6	6.0			
	Example 43	107	73.1	10.1			
	Example 44	66	73.8	6.6			
25	Example 45	340	78.1	26.8			
	Example 46	56	66.6	6.8			
	Comparative	104	83.1	10.0			
30	Example 21						
	Comparative	104	84.1	10.0			
	Example 22						
35	Comparative	103	83.1	10.1			
	Example 23						

Table 14 (continued)

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	r				
n	Properties of black magnetic composite				
Examples	particles				
and	Fluidity	Blackness	Carbon black		
Comparative	index	(L* value)	desorption		
Examples	(-)	(-)	percentage		
			(%)		
Example 39	52	16.0 ·	7.1		
Example 40	51	16.3	6.5		
Example 41	53	17.2	5.9		
Example 42	52	17.6	5.3		
Example 43	54	16.0	4.3		
Example 44	54	16.0	3.4		
Example 45	52	17.4	3.8		
Example 46	53	17.4	4.3		
Comparative	37	20.8	69.2		
Example 21					
Comparative	37	21.0	31.6		
Example 22					
Comparative	38	20.6	50.8		
Example 23					

Table 15

. 5					
3				ack magnetic to	ner
	Examples	Black magnetic		Resin	
	and	_composite p	articles		
	Comparative	Kind	Amount	Kind	Amount
10	Examples		blended		blended
			(part by		(part
			weight)		by
	Francis 42	D	45		weight)
15	Example 47	Example 23	45	Styrene-acryl	. 55
				copolymer	
	Example 48	Example 24	45	resin	
	Evaultie 40	Example 24	45	Styrene-acryl	55
20				copolymer resin	
	Example 49	Example 25	40	Styrene-acryl	60
		mampre 25	30	copolymer	60
				resin	
25	Example 50	Example 26	50	Styrene-acryl	50
20				copolymer	
				resin	
	Example 51	Example 27	45	Styrene-acryl	55
				copolymer	
30				resin	
	Example 52	Example 28	40	Styrene-acryl	60
				copolymer	
				resin	
35	Example 53	Example 29	50	Styrene-acryl	50
	;			copolymer	
	Francis 54	1 - 20		resin	
	Example 54	Example 30	50	Styrene-acryl	50
40				copolymer resin	
	Comparative	Comparative	45		55
	Example 24	Example 15	45	Styrene-acryl copolymer	22
		manupic 13		resin	
45	Comparative	Comparative	45	Styrene-acryl	55
	Example 25	Example 16		copolymer	33
				resin	[
	Comparative	Comparative	45	Styrene-acryl	55
<i>50</i>	Example 26	Example 17		copolymer	
55	_	/	İ	resin	

Table 15 (continued)

•				

Examples	Prope	Properties of black magnetic toner			
and	Average	Dispers-	Fluidity	Volume	
Comparative	particle	ibility	index	resistivity	
Examples	size	(-)	(-)	(Ω•cm)	
	(µm)		•		
Example 47	9.9	5	75	8.9x10 ¹³	
Example 48	10.0	5	81	1.8x10 ¹⁴	
Example 49	10.6	4	75	7.6x10 ¹³	
Example 50	10.5	5	78	7.1x10 ¹³	
Example 51	9.6	5	79	5.9x10 ¹³	
Example 52	9.9	5	83	3.1x10 ¹⁴	
Example 53	10.0	5	76	1.9x10 ¹⁴	
Example 54	10.8	5	83	1.5x10 ¹⁴	
Comparative Example 24	10.6	2	56	1.8x10 ¹²	
Comparative Example 25	10.5	2	58	2.1x10 ¹²	
Comparative Example 26	10.4	2.	56	2.1x10 ¹²	

Table 15 (continued)

Examples	Properties of black magnetic toner			
and	Ma	agnetic propert:	ies	
Comparative	Coercive	Saturation m	magnetization	
Examples	force	(10 k0e)	(1 k0e)	
	(Oe)	(emu/g)	(emu/g)	
Example 47	97	36.6	27.4	
Example 48	60	33.4	26.0	
Example 49	321	31.4	23.5	
Example 50	57	32.3	22.8	
Example 51	104	32.5	23.5	
. Example 52	67	29.8	22.6	
Example 53	328	33.6	23.6	
Example 54	52	31.7	24.1	
Comparative	101	37.2	27.2	
Example 24				
Comparative	102	38.0	29.3	
Example 25				
Comparative	102	36.5	28.5	
Example 26				

Table 15 (continued)

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Examples	Properties of black magnetic toner			
and	Magnetic	properties	Blackness	
Comparative	Residual ma	agnetization	(L* value)	
Examples	(10 kOe)	(1 k0e)	(-)	
	(emu/g)	(emu/g)		
Example 47	5.8	4.3	18.5	
Example 48	4.1	2.9	17.9	
Example 49	11.1	8.4	19.2	
Example 50	4.4	3.2	19.3	
Example 51	5.3	4.0	17.6	
. Example 52	3.7	2.8	18.2	
Example 53	10.1	7.2	18.0	
Example 54	4.7	3.1	19.1	
Comparative	5.1	4.0	22.2	
Example 24				
Comparative	5.1	4.0	23.6	
Example 25				
Comparative	5.0	4.1	23.0	
Example 26				

Table 16

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	Production of black magnetic toner			
Examples	Black magnetic Resin			
and	composite p	articles		
Comparative	Kind	Amount	Kind	Amount
Examples		blended		blended
		(part by		(part
		weight)		by
		·		weight)
Example 55	Example 31	45	Styrene-acryl	55
			copolymer	
			resin	
Example 56	Example 32	45	Styrene-acryl	55
			copolymer	
			resin	
Example 57	Example 33	40	Styrene-acryl	60
_	_		copolymer	
			resin	
Example 58	Example 34	50	Styrene-acryl	50
_	-		copolymer	
			resin	
Example 59	Example 35	45	Styrene-acryl	55
			copolymer	
			resin	
Example 60	Example 36	40	Styrene-acryl	60
	_		copolymer	
			resin	
Example 61	Example 37	50	Styrene-acryl	50
	_		copolymer	·
			resin	
Example 62	Example 38	50	Styrene-acryl	50
	·		copolymer	
			resin	
Comparative	Comparative	45	Styrene-acryl	55
Example 27	Example 18	_	copolymer	
-	-		resin	
Comparative	Comparative	45	Styrene-acryl	55
Example 28	Example 19		copolymer	
·	•		resin	
Comparative	Comparative	45	Styrene-acryl	55
Example 29	Example 20		copolymer	
			resin	
L				

Table 16 (continued)

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Examples	Prope	erties of bl	ack magneti	c toner
and	Average	Dispers-	Fluidity	Volume
Comparative	particle	ibility	index	resistivity
Examples	size	(-)	(∸)	(Ω•cm)
	(µm)	;		
Example 55	10.0	5	76	9.2×10 ¹³
Example 56	10.0	5	80	2.5x10 ¹⁴
Example 57	10.1	4	76	6.5x10 ¹³
Example 58	9.9	5	81	7.8x10 ¹³
Example 59	10.1	5	80	7.3x10 ¹³
Example 60	9.8	5	85	3.2x10 ¹⁴
Example 61	10.2	5	75	2.6x10 ¹⁴
Example 62	10.0	5	83	1.4x10 ¹⁴
Comparative Example 27	10.2	2	60	1.8x10 ¹²
Comparative Example 28	10.4	2	59	3.1x10 ¹²
Comparative Example 29	10.2	2	60	3.4x10 ¹²

Table 16 (continued)

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15	
20	
25	
30	
35	

	· · · · · · · · · · · · · · · · · · ·			
Examples	Properties of black magnetic toner			
and	Ma	agnetic propert	ies	
Comparative	Coercive	Saturation n	magnetization	
Examples	force	(10 kOe) .	(1 kOe)	
	(0e)	(emu/g)	(emu/g)	
Example 55	96	36.5	27.4	
Example 56	61	33.8	26.5	
Example 57	321	34.4	23.2	
Example 58	58	32.6	22.4	
Example 59	102	32.0	23.4	
Example 60	66	29.4	22.1	
Example 61	318	33.4	25.6	
Example 62	51	31.6	23.4	
Comparative	101	38.2	27.8	
Example 27				
Comparative	102	38.2	29.3	
Example 28				
Comparative	100	36.7	28.5	
Example 29				

Table 16 (continued)

5					
	Examples	Propertie	Properties of black magne		
	and	Magnetic properties		Blackness	
10	Comparative	Residual ma	Residual magnetization		
	Examples	(10 kOe)	(1 kOe) .	(-)	
15		(emu/g)	(emu/g)		
	Example 55	5.8	4.3	18.4	
	Example 56	4.1	2.8	17.9	
20	Example 57	11.4	8.6	19.3	
	Example 58	4.5	. 3.3	18.8	
	Example 59	5.3	4.1	17.2	
25	Example 60	3.7	2.8	17.9	
	Example 61	8.6	6.4	18.3	
	Example 62	4.3	3.1	18.6	
30	Comparative	5.9	4.6	23.3	
	Example 27				
35	Comparative	5.3	4.1	22.8	
	Example 28				
	Comparative	5.0	3.9	22.2	
40	Example 29				

Table 17

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	Production of black magnetic toner			
Examples	Black magnetic		Resin	
and	_partic	les		
Comparative	Kind	Amount	Kind	Amount
Examples		blended		blended
}		(part by		(part
		weight)		by
				weight)
Example 63	Example 39	45	Styrene-acryl	55
_	_		copolymer	
1			resin	
Example 64	Example 40	45	Styrene-acryl	55
_	_		copolymer	
			resin	
Example 65	Example 41	40	Styrene-acryl	60
			copolymer	
			resin	
Example 66	Example 42	50	Styrene-acryl	50
			copolymer	30
			resin	
Example 67	Example 43	45	Styrene-acryl	55
			copolymer	33
1			resin	
Example 68	Example 44	40	Styrene-acryl	60
	-		copolymer	
1			resin	
Example 69	Example 45	50	Styrene-acryl	50
			copolymer	
			resin	
Example 70	Example 46	50	Styrene-acryl	50
			copolymer	
1			resin	·
Comparative	Comparative	45	Styrene-acryl	55
Example 30	Example 21		copolymer	33
			resin	
Comparative	Comparative	45	Styrene-acryl	55
Example 31	Example 22		copolymer	,,
		ľ	resin	
Comparative	Comparative	45	Styrene-acryl	55
Example 32	Example 23	±2	copolymer	25
Transpie 12	TVERIFITE 53		resin	
<u> </u>		-	rearm	

Table 17 (continued)

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Examples	Properties of black magnetic toner			
and	Average	Dispers-	Fluidity	Volume
Comparative	particle	ibility	index	resistivity
Examples	size	(-)	(-)	(Ω•cm)
	(µm)			
Example 63	10.1	5	75	8.6x10 ¹³
Example 64	9.8	5	78	2.1x10 ¹⁴
Example 65	10.2	4	72	6.5x10 ¹³
Example 66	9.9	5	80	7.3x10 ¹³
Example 67	10.3	5	80	7.1x10 ¹³
Example 68	10.0	5	82	3.2x10 ¹⁴
Example 69	9.6	5	79	1.6x10 ¹⁴
Example 70	10.0	5	83	2.1x10 ¹⁴
Comparative	9.8	2	59	1.2x10 ¹²
Example 30				
Comparative	9.9	2	57	1.4x10 ¹²
Example 31				
Comparative	10.0	2	57	3.2x10 ¹²
Example 32				

Table 17 (continued)

5	

Examples	Properties of black magnetic toner		
and	Magnetic properties		
Comparative	Coercive	Saturation m	nagnetization
Examples	force	(10 kOe)	(1 k0e)
	(0e)	(emu/g)	(emu/g)
Example 63	98	36.9	27.3
Example 64	62	33.8	26.2
Example 65	308	34.9	23.1
Example 66	58	32.6	22.6
Example 67	101	32.6	23.1
Example 68	64	32.6	22.6
Example 69	313	33.2	24.2
Example 70	56	32.1	23.1
Comparative	102	38.6	27.3
Example 30			
Comparative	103	37.9	25.6
Example 31			
Comparative	101	37.1	28.3
Example 32			

Table 17 (continued)

Examples	Properties of black magnetic toner		
and	Magnetic	properties	Blackness
Comparative	Residual ma	gnetization	(L* value)
Examples	(10 kOe)	(1 k0e)	(-)
	(emu/g)	(emu/g)	
Example 63	5.6	4.2	18.6
Example 64	3.6	2.8	17.8
Example 65	12.1	8.4	19.0
Example 66	3.6	3.1	18.6
Example 67	5.1	3.9	17.4
. Example 68	3.8	2.5	18.0
Example 69	10.6	5.2	18.3
Example 70	4.3	3.4	18.9
Comparative	5.8	4.3	23.2
Example 30			
Comparative	5.3	4.2	23.5
Example 31		!	
Comparative	5.0	4.0	23.1
Example 32	ļ		

Table 18

. 5				
	Examples	Kind of	Production of black	magnetic
	and	particles	composite partic	les
10	Comparative	to be	Coating step wi	
,,	Examples	treated	fluoroalkylsilane co	
			Fluoroalkylsilane co	Amount
				added
15				(part by
				weight)
	Example 71	Core	Tridecafluorooctyl	2.0
20		particles 1	trimethoxysilane	
	Example 72	Core	Heptadecafluorodecyl	4.0
		particles 2	trimethoxysilane	
	Example 73	Core	Trifluoropropyl	3.0
25		particles 3	trimethoxysilane	
	Example 74	Core	Tridecafluorooctyl	1.0
		particles 4	trimethoxysilane	
30	Example 75	Core	Tridecafluorooctyl	6.0
		particles 5	trimethoxysilane	
	Example 76	Core	Heptadecafluorodecyl	4.0
35		particles 6	trimethoxysilane	
	Example 77	Core	Trifluoropropyl	0.5
		particles 7	trimethoxysilane	
40	Example 78	Core	Tridecafluorooctyl	1.5
		particles 8	trimethoxysilane	
	Comparative	Core	Tridecafluorooctyl	2.0
45	Example 33	particles 1	trimethoxysilane	
~	Comparative	Core	Tridecafluorooctyl	3.0
	Example 34	particles 1	trimethoxysilane	
	Comparative	Core	Tridecafluorooctyl	0.005
50	Example 35	particles 1	trimethoxysilane	

Table 18 (continued)

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Example		Production	of black magnet:	ic composite	
Example	_	Production of black magnetic composite			
	S	particles			
and		Coating step w	ith fluoroalkyl:	silane compound	
Comparati	ve	Edge runne	r treatment	Coating amount	
Example	s	Linear load	Time	(calculated	
		(Kg/cm)	(min)	as Si)	
		****	•	(wt. %)	
Example '	71	60	30	0.13	
Example '	72	45	25	0.20	
Example '	73	30	40	0.47	
Example '	74	75	45	0.07	
Example 7	/5	60	30	0.39	
Example 7	6	60	20	0.21	
Example 7	7	30	45	0.08	
Example 7	8	30	60	0.10	
Comparati	ve	60	30	0.13	
Example 3	3				
Comparati	vе	60	30	0.21	
Example 3	4				
Comparation	re	60	30	3.0×10 ⁻⁴	
Example 3	5				

Table 18 (continued)

5			•	
		Production of black magnetic composite		
	Examples	particles		
10	and	Adhering step of carbo	on black fine particles	
	Comparative	Carbon black	fine particles	
	Examples	Kind	Amount added	
15			(part by weight)	
15	Example 71	A	. 8.0	
	Example 72	A	6.0	
20	Example 73	В	5.0	
20	Example 74	С	13.0	
	Example 75	A	10.0	
25	Example 76	A	10.0	
	Example 77	В	18.0	
	Example 78	С	16.0	
30	Comparative	-	- 1. 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	
	Example 33			
	Comparative	A	0.01	
35	Example 34			
	Comparative	В	5.0	
	Example 35			

Table 18 (continued)

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	Production of black magnetic composite		
Examples	particles		
and	Adhering step	of carbon black	fine particles
Comparative	Edge runne:	r treatment	Amount adhered
Examples	Linear load	Time	(calculated
1	(Kg/cm)	(min)	as C)
		•	(wt. %)
Example 71	60	30	7.41
Example 72	30	45	5.69
Example 73	60	30	4.79
Example 74	30	45	11.53
Example 75	60	25	9.00
Example 76	30	. 60	9.10
Example 77	60	45	15.17
Example 78	30	45	13.82
Comparative	_	_	_
Example 33			
Comparative	60	30	0.01
Example 34			
Comparative	60	30	4.75
Example 35			

Table 19

				·
Examples	Properties of black magnetic composite particles			
_				····
and	Average	Aspect	Geometrical	BET
Comparative	particle	ratio	standard	specific
Examples	size	(–)	deviation	surface
	(µm)		(-)	area
			·	(m^2/g)
Example 71	0.29	-	1.53	6.3
Example 72	0.23	_	1.33	12.8
Example 73	0.40	8.1:1	1.52	26.8
Example 74	0.23	-	1.43	14.6
Example 75	0.29	_	1.53	15.3
Example 76	0.23	_	1.33	14.8
Example 77	0.40	8.1:1	1.51	28.8
Example 78	0.23	-	1.43	13.4
Comparative	0.28	-	1.52	10.0
Example 33		•		
Comparative	0.29	-	1.52	9.8
Example 34				:
Comparative	0.29	···	1.52	13.6
Example 35				

Table 19 (continued)

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Examples and Comparative Examples Coercive Examples Coercive Examples Coercive Example 71 108 81.5 11.7		Properties of black magnetic composite			
Comparative Examples Coercive force (Oe) Saturation magnetization (10 kOe) (emu/g) Residual magnetization (10 kOe) (emu/g) Example 71 108 81.5 11.7 Example 72 68 72.4 6.5 Example 73 343 74.6 25.3 Example 74 58 64.3 6.1 Example 75 107 73.2 10.0 Example 76 67 73.6 6.5 Example 77 340 78.1 26.3 Example 78 57 63.2 6.5 Comparative 103 83.3 9.9 Example 34 10.2 9.8 Comparative 103 83.3 10.2	Examples	particles			
Examples force (Oe) magnetization (10 kOe) (emu/g) magnetization (10 kOe) (emu/g) Example 71 108 81.5 11.7 Example 72 68 72.4 6.5 Example 73 343 74.6 25.3 Example 74 58 64.3 6.1 Example 75 107 73.2 10.0 Example 76 67 73.6 6.5 Example 77 340 78.1 26.3 Example 78 57 63.2 6.5 Comparative 103 83.3 9.9 Example 33 Comparative 103 84.1 9.8 Example 34 Comparative 103 83.3 10.2		Magnetic properties			
(Oe) (10 kOe) (emu/g) (10 kOe) (emu/g) Example 71 108 81.5 11.7 Example 72 68 72.4 6.5 Example 73 343 74.6 25.3 Example 74 58 64.3 6.1 Example 75 107 73.2 10.0 Example 76 67 73.6 6.5 Example 77 340 78.1 26.3 Example 78 57 63.2 6.5 Comparative 103 83.3 9.9 Example 33 84.1 9.8 Example 34 20 83.3 10.2	_	Coercive	Saturation	Residual	
Example 71 108 81.5 11.7 Example 72 68 72.4 6.5 Example 73 343 74.6 25.3 Example 74 58 64.3 6.1 Example 75 107 73.2 10.0 Example 76 67 73.6 6.5 Example 77 340 78.1 26.3 Example 78 57 63.2 6.5 Comparative 103 83.3 9.9 Example 33 84.1 9.8 Example 34 103 83.3 10.2	Examples	force	magnetization	magnetization	
Example 71 108 81.5 11.7 Example 72 68 72.4 6.5 Example 73 343 74.6 25.3 Example 74 58 64.3 6.1 Example 75 107 73.2 10.0 Example 76 67 73.6 6.5 Example 77 340 78.1 26.3 Example 78 57 63.2 6.5 Comparative 103 83.3 9.9 Example 34 Comparative 103 84.1 9.8 Example 34 Comparative 103 83.3 10.2		(0e)	(10 kOe)	(10 kOe)	
Example 72 68 72.4 6.5 Example 73 343 74.6 25.3 Example 74 58 64.3 6.1 Example 75 107 73.2 10.0 Example 76 67 73.6 6.5 Example 77 340 78.1 26.3 Example 78 57 63.2 6.5 Comparative 103 83.3 9.9 Example 33 Comparative 103 84.1 9.8 Example 34 Comparative 103 83.3 10.2					
Example 73 343 74.6 25.3 Example 74 58 64.3 6.1 Example 75 107 73.2 10.0 Example 76 67 73.6 6.5 Example 77 340 78.1 26.3 Example 78 57 63.2 6.5 Comparative 103 83.3 9.9 Example 33 Comparative 103 84.1 9.8 Example 34 Comparative 103 83.3 10.2					
Example 74 58 64.3 6.1 Example 75 107 73.2 10.0 Example 76 67 73.6 6.5 Example 77 340 78.1 26.3 Example 78 57 63.2 6.5 Comparative 103 83.3 9.9 Example 33 Comparative 103 84.1 9.8 Example 34 Comparative 103 83.3 10.2	Example 72	68	72.4	6.5	
Example 75 107 73.2 10.0 Example 76 67 73.6 6.5 Example 77 340 78.1 26.3 Example 78 57 63.2 6.5 Comparative 103 83.3 9.9 Example 33 Comparative 103 84.1 9.8 Example 34 Comparative 103 83.3 10.2	Example 73	343	74.6	25.3	
Example 76 67 73.6 6.5 Example 77 340 78.1 26.3 Example 78 57 63.2 6.5 Comparative 103 83.3 9.9 Example 33 Comparative 103 84.1 9.8 Example 34 Comparative 103 83.3 10.2	Example 74	58	64.3	6.1	
Example 77 340 78.1 26.3 Example 78 57 63.2 6.5 Comparative 103 83.3 9.9 Example 33 Comparative 103 84.1 9.8 Example 34 Comparative 103 83.3 10.2	Example 75	107	73.2	10.0	
Example 78 57 63.2 6.5 Comparative 103 83.3 9.9 Example 33 Comparative 103 84.1 9.8 Example 34 Comparative 103 83.3 10.2	Example 76	67	73.6	6.5	
Comparative 103 83.3 9.9 Example 33 103 84.1 9.8 Example 34 20 103 83.3 10.2	Example 77	340	78.1	26.3	
Example 33 Comparative 103 84.1 9.8 Example 34 Comparative 103 83.3 10.2	Example 78	57	63.2	6.5	
Comparative 103 84.1 9.8 Example 34 Sample 34 10.2	Comparative	103	83.3	9.9	
Example 34	Example 33				
Comparative 103 83.3 10.2	Comparative	103	84.1	9.8	
	Example 34				
Example 35	Comparative	103	83.3	10.2	
	Example 35				

Table 19 (continued)

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	Properties of black magnetic composite		
Examples	particles		
and	Fluidity	Blackness	Carbon black
Comparative	index	(L* value)	desorption
Examples	(-)	. (-)	percentage
			(융)
Example 71	47	16.4	6.1
Example 72	48	16.8	7.4
Example 73	50	17.3	8.2
Example 74	46	17.8	5.6
Example 75	51	16.5	4.3
Example 76	52	16.3	4.1
Example 77	53	17.6	3.8
Example 78	52	17.8	4.8
Comparative	38	20.6	79.1
Example 33			
Comparative	38	20.8	28.7
Example 34			
Comparative	37	20.3	53.4
Example 35			

Table 20

_					•
5		Product	Production of black magnetic toner		
	Examples	Black mag	metic	Resin	
	and	partic	les		
i	Comparative	Kind	Amount	Kind	Amount
10	Examples		blended		blended
			(part by		(part
		:	weight)		by
					weight)
15	Example 79	Example 71	45	Styrene-acryl	55
ĺ				copolymer	
				resin	
	Example 80	Example 72	45	Styrene-acryl	55
20		. •		copolymer	
20				resin	
	Example 81	Example 73	40	Styrene-acryl	60
		_		copolymer	
				resin	50
25	Example 82	Example 74	50	Styrene-acryl	50
				copolymer resin	
			45	Styrene-acryl	55
	Example 83	Example 75	45	copolymer	33
30				resin	
	Example 84		40	Styrene-acryl	60
	EXAMPLE 04	Example 76		copolymer	
				resin	
35	Example 85	5 1 27	50	Styrene-acryl	50
		Example 77		copolymer	
				resin	
	Example 86	Example 78	50	Styrene-acryl	50
	•	example 10		copolymer	
40				resin	
	Comparative	Comparative	45	Styrene-acryl	55
	Example 36	_		copolymer	
		Example 33		resin	
45	Comparative	Comparative	45	Styrene-acryl	55
	Example 37			copolymer	
		Example 34		resin	
	Comparative	Comparative	45	Styrene-acryl	55
50	Example 38			copolymer	
		Example 35	<u></u>	resin	

Table 20 (continued)

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Examples	Properties of black magnetic toner			
and	Average	Dispers-	Fluidity	Volume
Comparative	particle	ibility	index	resistivity
Examples	size	(-)	(∸)	(Ω•cm)
	(µm)			
Example 79	10.0	5	76	8.1x10 ¹³
Example 80	9.8	4	81	2.1x10 ¹⁴
Example 81	10.1	4	73	6.5x10 ¹³
Example 82	10.3	5	82	9.2x10 ¹³
Example 83	10.2	5	85	5.4x10 ¹³
Example 84	10.0	5	86	3.6x10 ¹⁴
Example 85	9.9	5	83	2.6x10 ¹⁴
Example 86	9.8	5	84	3.8x10 ¹⁴
Comparative	10.1	2	57	1.3x10 ¹²
Example 36				`
Comparative	10.3	2	56	2.4x10 ¹²
Example 37				
Comparative	10.1	2	56	6.8x10 ¹²
Example 38				

Table 20 (continued)

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P	Properties of black magnetic toner			
Examples	Properties	s of black magn	etic toner	
and	Ma	gnetic properti	.es	
Comparative	Coercive	Saturation m	agnetization	
Examples	force	(10 kOe)	(1 k0e)	
	(Oe)	(emu/g)	(emu/g)	
Example 79	100	36.9	27.0	
Example 80	63	34.8	26.4	
Example 81	315	34.3	22.9	
Example 82	58	31.6	22.6	
Example 83	101	32.1	23.3	
Example 84	61	32.8	22.2	
Example 85	318	32.6	24.1	
Example 86	55	32.1	23.0	
Comparative	101	38.4	27.3	
Example 36				
Comparative	100	38.0	25.3	
Example 37				
Comparative	101	37.1	27.2	
Example 38				

Table 20 (continued)

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Examples	Properties of black magnetic toner			
and	Magnetic	properties	Blackness	
Comparative	Residual ma	agnetization	(L* value)	
Examples	(10 kOe)	(1 kOe)	(-)	
	(emu/g)	(emu/g)		
Example 79	5.7	4.1	18.5	
Example 80	3.7	2.8	17.6	
Example 81	12.3	8.3	18.8	
Example 82	3.8	3.0	18.6	
Example 83	5.6	3.4	17.2	
Example 84	3.6	2.1	18.3	
Example 85	9.8	5.3	17.3	
Example 86	3.7	3.2	17.3	
Comparative	5.4	4.3	23.4	
Example 36				
Comparative	5.3	4.5	23.2	
Example 37				
Comparative	5.0	4.0	22.2	
Example 38				

45 Claims

- 1. Black magnetic composite particles for a black magnetic toner, comprising:
 - (a) magnetic iron oxide particles having an average particle diameter of from 0.055 to 0.95 μ m;
 - (b) a coating on the surface of said magnetic iron oxide particles, comprising at least one organosilicon compound selected from:
 - (1) organosilane compounds obtainable by drying or heat-treating alkoxysilane compounds,
 - (2) polysiloxanes or modified polysiloxanes, and
 - (3) fluoroalkyl organosilane compounds obtainable by drying or heat-treating fluoroalkylsilane compounds; and
 - (c) carbon black fine particles adhered on at least a part of said coating, which hav a particl siz of from

0.002 to $0.05 \,\mu m$ and which are present in an amount of from 1 to 25 parts by weight per 100 parts by weight of said magn tic iron oxide particles.

- 2. Particles according to claim 1, wherein said magnetic iron oxide particles have a coat on at least a part of th surface thereof, which comprises at least one hydroxide or oxide selected from hydroxides of aluminum, oxides of aluminum, hydroxides of silicon and oxides of silicon in an amount of from 0.01 to 50 % by weight, calculated as Al and/or SiO₂, based on the total weight of the magnetic iron oxide particles.
 - 3. Particles according to claim 1 or 2, wherein said alkoxysilane compound is represented by the general formula (I):

$$R^{1}_{a}SiX_{4-a}$$
 (I)

wherein R¹ is C_6H_{5-} , $(CH_3)_2CHCH_2$ - or $n-C_bH_{2b+1}$ - (wherein b is an integer of from 1 to 18); X is CH_3O - or C_2H_5O -; and a is an integer of from 0 to 3.

- 4. Particles according to claim 3, wherein said alkoxysilane compound is methyl triethoxysilane, dimethyl diethoxysilane, tetraethoxysilane, phenyl triethoxysilane, diphenyl diethoxysilane, methyl trimethoxysilane, dimethyl dimethoxysilane, tetramethoxysilane, phenyl trimethoxysilane, diphenyl dimethoxysilane, isobutyl trimethoxysilane or decyl trimethoxysilane.
- Particles according to any one of the preceding claims, wherein said polysiloxanes are represented by the general formula (II):

wherein R² is H- or CH₃-, and d is an integer of 15 to 450.

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- Particles according to claim 5, wherein said polysiloxanes have methyl hydrogen siloxane units.
 - 7. Particles according to any one of the preceding claims, wherein said modified polysiloxanes are selected from:
 - (A) polysiloxanes modified with at least one polyether and/or polyester and/or epoxy compound, and
 (B) polysiloxanes whose molecular terminal is modified with at least one group selected from carboxylic acid, alcohol and hydroxyl groups.
 - 8. Particles according to claim-7, wherein said polysiloxanes (A) are represented by the general formula (III), (IV) or (V):

wherein R^3 is -(-CH₂-)_h-; R^4 is -(-CH₂-)_i-CH₃; R^5 is -OH, -COOH, -CH=CH₂, -C(CH₃)=CH₂ or -(-CH₂-)_j-CH₃; R^6 is -(-CH₂-)_k-CH₃; R^6 and R^6 and R^6 is -(-CH₂-)_k-CH₃; R^6 is an integer of from 1 to 15; i, j and k are integers of from 0 to 15; e is an integer of from

1 to 50; and f is an integer of from 1 to 300;

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wherein R⁷, R⁸ and R⁹ are -(-CH₂-)_q- and may be the same or different; R¹⁰ is -OH, -COOH, -CH=CH₂, -C(CH₃) =CH₂ or -(-CH₂-)_r-CH₃; R¹¹ is -(-CH₂-)_s-CH₃; n and q are integers of from 1 to 15; r and s are integers of from 0 to 15; e' is an integer of from 1 to 50; and f' is an integer of from 1 to 300; or

wherein R^{12} is -(- CH_2 -)_v-; v is an integer of from 1 to 15; t is an integer of from 1 to 50; and u is an integer of from 1 to 300.

9. Particles according to claim 7 or 8, wherein said polysiloxanes (B) are represented by the general formula (VI):

wherein R¹³ and R¹⁴ are -OH, R¹⁶OH or R¹⁷COOH and may be the same or different; R¹⁵ is -CH₃ or -C₆H₅; R¹⁶ and R¹⁷ are -(-CH₂-)_y-; y is an integer of from 1 to 15; w is an integer of from 1 to 200; and x is an integer of from 0 to 100.

 Particles according to any one of the preceding claims, wherein said fluoroalkylsilane compounds are represented by the general formula (VII):

$$CF_3(CF_2)_z CH_2 CH_2(R^{18})_a SiX_{4-a}$$
 (VII)

wherein R¹⁸ is CH₃-, C₂H₅-, CH₃O- or C₂H₅O-; X is CH₃O- or C₂H₅O-; and z is an integer of from 0 to 15; and a' is an integer of from 0 to 3.

- 11. Particles according to any one of the preceding claims, wherein the amount of said coating organosilicon compound (s) is from 0.02 to 5.0 % by weight, calculated as Si, based on the total weight of the organosilicon compound(s) and said magnetic iron oxide particles.
- 12. Particles according to any one of the preceding claims, which have an average particle diameter of from 0.06 to

1.0 µm

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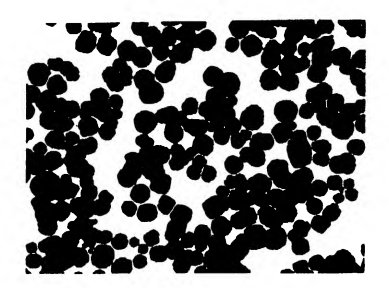
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- 13. Particles according to any one of the pr ceding claims, which hav a geometrical standard deviation of particle siz s of from 1.01 to 2.0.
- 14. Particles according to any one of the preceding claims, which have a BET specific surface area value of from 1 to 200 m²/g.
- 15. Particles according to any one of the preceding claims, which have a flowability index of from 45 to 80.
- 16. Particles according to any one of the preceding claims, which have a blackness (L* value) of from 15 to 20.
- 17. Black magnetic toner comprising said black magnetic composite particles according to any one of the preceding claims.
- **18.** Toner according to claim 17, which comprises from 50 to 900 parts by weight of a binder resin per 100 parts by weight of said black magnetic composite particles.
- 19. Toner according to claim 17 or 18, which has an average particle size of from 3 to 15 μm .
- 20. Toner according to any one of claims 17 to 19, which has a flowability index of from 70 to 100.
- 21. Toner according to any one of claims 17 to 20, which has a blackness (L* value) of from 15 to 20.
- 25 22. Toner according to any one of claims 17 to 21, which has a volume resistivity of from 1.0 x 10^{13} to 1.0 x 10^{15} Ω .
 - 23. A method for production of a black magnetic toner, which method comprises mixing black magnetic composite particles according to any one of claims 1 to 16 with a binder resin and processing the resulting mixture thereby to form the toner.

FIG.1



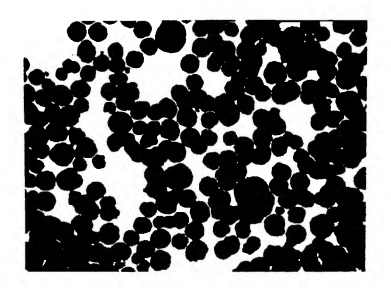
(×20000)

FIG.2



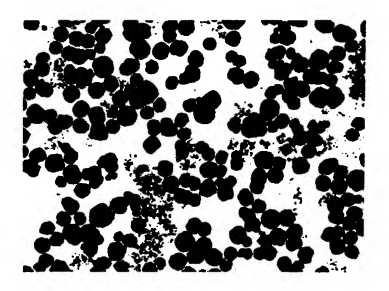
(×20000)

FIG.3



(×20000)

FIG.4



(×20000)